as a sweetening agent methyl and propylparabous as preservatives, a dye and flavoring, such as cherry or orange flavor. Of course, any material used in preparing any dosage unit form should be pharmaceutically pure and substantially non-toxic in the amounts employed. In addition, the active compounds may be incorporated into sustained-release preparation and formulations.

Typically, these formulations may contain at least about 0.1% of the active compound or more, although the percentage of the active ingredient(s) may, of course, be varied and may conveniently be between about 1 or 2% and about 60% or 70% or more of the weight or volume of the total formulation. Naturally, the amount of active compound(s) in each therapeutically useful composition may be prepared is such a way that a suitable dosage will be obtained in any given unit dose of the compound. Factors such as solubility, bioavailability, biological half-life, route of administration, product shelf life, as well as other pharmacological considerations will be contemplated by one skilled in the art of preparing such pharmaceutical formulations, and as such, a variety of dosages and treatment regimens may be desirable.

For oral administration the compositions of the present invention may alternatively be incorporated with one or more excipients in the form of a mouthwash, dentifrice, buccal tablet, oral agray, or sublingual orally-administered formulation. For example, a mouthwash may be prepared incorporating the active ingredient in the required amount in an appropriate solvent, such as a sodium borate solution (Dobell's Solution). Alternatively, the active ingredient may be incorporated into an oral solution such as one containing sodium borate, glycerin and potassium bicarbonate, or dispersed in a dentifrice, or added in a therapeutically-effective amount to a composition that may include water, binders, abrasives, flavoring agents, feaming agents, and humectants. Alternatively the compositions may be fashioned into a tablet or solution form that may be placed under the tongue or otherwise dissolved in the mouth.

2. Injectable delivery

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In certain circumstances it will be desirable to deliver the pharmaceutical compositions disclosed herein parenterally, intravenously, intramuscularly, or even intraperitoneally as described in, e.g., U.S. Patent Nos. 5,543,158; 5,641,515; and 5,399,363. Solutions of the active compounds as free base or pharmacologically

acceptable salts may be prepared in water suitably mixed with a surfactant, such as hydroxypropylecllulose. Dispersions may also be prepared in glycerol, liquid polyethylene glycols, and mixtures thereof and in oils. Under ordinary conditions of storage and use, these preparations contain a preservative to prevent the growth of microorganisms.

The pharmaceotical forms suitable for injectable use include sterile aqueous solutions or dispersions and sterile powders for the extemporaneous preparation of sterile injectable solutions or dispersions (U.S. Patent No. 5,466,468). In all cases the form must be sterile and must be fluid to the extent that easy syringsbility exists. It must be stable under the conditions of manufacture and storage and must be preserved against the contaminating action of microorganisms, such as bacteria and fangi. The carrier can be a solvent or dispersion medium containing, for example, water, ethanol, polyol (e.g., glycerol, propylene glycol, and liquid polyethylene glycol, and the like), suitable mixtures thereof, and/or vegetable oils. Proper fluidity may be maintained, for example, by the use of a coating, such as legithin, by the maintenance of the required particle size in the case of dispersion and by the use of surfactants. The prevention of the action of microorganisms can be facilitated by various antibacterial and antifungal agents, for example, parabens, chlorobatanol, phenol, sorbic acid, thimerosal, and the like. In many cases, it will be preferable to include isotonic agents, for example, sugars or sodium chloride. Prolonged absorption of the injectable compositions can be brought about by the use in the compositions of agents delaying absorption, for example, aluminum monostearate and gelatin.

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For parenteral administration in an aqueous solution, for example, the solution should be suitably buffered if necessary and the liquid diluent first rendered isotenic with sufficient saline or glucose. These particular aqueous solutions are especially suitable for infravenous, intramuscular, subcutaneous and intraperitoneal administration. In this consection, a sterile aqueous medium that can be employed will be known to those of skill in the art in high of the present disclosure. For example, one dosage may be dissolved in 1 ml of isotonic NaCl solution and either added to 1000 ml of hypodermoelysis fluid or injected at the proposed site of infusion (see, e.g., Remington Pharmaceutical Sciences 15th Edition, pages 1035-1038 and 1570-1580). Some variation in dosage will necessarily occur depending on the condition of the subject being treated.

The person responsible for administration will, in any event, determine the appropriate dose for the individual subject. Moreover, for human administration, preparations should meet sterility, pyrogenicity, and the general safety and purity standards as required by FDA Office of Biologics standards.

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Sterile injectable solutions are prepared by incorporating the active compounds in the required amount in the appropriate solvent with various of the other ingredients enumerated above, as required, followed by filtered sterilization. Generally, dispersions are prepared by incorporating the various sterilized active ingredients into a sterile vehicle which contains the basic dispersion medium and the required other ingredients from those enumerated above. In the case of sterile powders for the preparation of sterile injectable solutions, the preferred methods of preparation are vacuum-drying and freeze-drying techniques which yield a powder of the active ingredient plus any additional desired ingredient from a previously sterile-filtered solution thereof.

The compositions disclosed herein may be formulated in a neutral or salt form. Pharmacoutically-acceptable salts, include the acid addition salts (formed with the free amine groups of the protein) and which are formed with inorganic acids such as, for example, hydrochloric or phosphotic acids, or such organic acids as acetic, axalic, tartaric, mandelic, and the like. Salts formed with the free carboxyl groups can also be derived from inorganic bases such as, for example, sodium, potassium, ammonium, calcium, or ferric hydroxides, and such organic bases as isopropylamine, trimethylamine, histidine, procaine and the like. Upon formulation, solutions will be administered in a manner compatible with the dosage formulation and in such amount as is therapeutically effective. The formulations are easily administered in a variety of dosage forms such as infectable solutions, drex-release capsules, and the like.

As used herein, "carrier" includes any and all solvents, dispersion media, vehicles, coatings, diluents, antibacterial and antifungal agents, isotonic and absorption delaying agents, buffers, carrier solutions, suspensions, colloids, and the like. The use of such media and agents for pharmaceutical active substances is well known in the art. Except insofar as any conventional media or agent is incompatible with the active ingredient, its use in the therapeutic compositions is contemplated. Supplementary active ingredients can also be incorporated into the compositions.

The phrase "pharmacoutically-acceptable" refers to molecular entities and compositions that do not produce an allergic or similar autoward reaction when administered to a human. The proparation of an aqueous composition that contains a protein as an active ingredient is well understood in the art. Typically, such compositions are prepared as injectables, either as liquid solutions or suspensions; solid forms suitable for solution in, or suspension in, liquid prior to injection can also be prepared. The preparation can also be emulsified.

3. Nasal delivery

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In certain erabodiments, the pharmaceutical compositions may be delivered by intranasal sprays, inhalation, and/or other aerosol delivery vehicles. Methods for delivering genes, nucleic acids, and peptide compositions directly to the lungs via nasal aerosol sprays has been described e.g., in U.S. Patent Nos. 5,756,353 and 5,804,212. Likewise, the delivery of drugs using intranasal microparticle resins (Takenaga et al., J Controlled Release 52:81-87 (1998)) and lysophosphatidyl-glycerol compounds (see, e.g., U.S. Patent No. 5,725,871) are also well-known in the pharmaceutical arts. Likewise, transmucosal drug delivery in the form of a polyeterafluoroetheylene support matrix is described in U.S. Patent No. 5,780,045.

4. Liposome-, nanocapsule-, and microparticle-mediated delivery

In certain embodiments, the inventors contemplate the use of liposomes, nanocapsules, microparticles, microspheres, lipid particles, vesicles, and the like, for the introduction of the compositions of the present invention into suitable host cells. In particular, the compositions of the present invention may be formulated for delivery either encapsulated in a lipid particle, a liposome, a vesicle, a nanosphere, or a nanoparticle or the like.

Such formulations may be preferred for the introduction of pharmaceutically-acceptable formulations of the polypeptides, fusion proteins and moleic acids disclosed herein. The formation and use of liposomes is generally known to those of skill in the art (see, e.g., Couvreur et al., FEBS Lett. 84(2):323-326 (1977); Couvreur (1988); Lasic, Trends Biotechnol. 16(7):307-321 (1998); which describes the use of liposomes and nanocapsules in the targeted antibiotic therapy for intracellular bacterial

infections and diseases). Recently, liposomes were developed with improved serum stability and circulation half-times (Gabizon and Papahadjopoulos, Proc Natl Acad Sci U S.A. 85(18):6949-6953 (1988); Allen and Choun (1987); U.S. Patent No. 5,741,516). Further, various methods of liposome and liposome like preparations as potential drug carriers have been reviewed (Takakura, Nippon Rinsho 56(3):691-695 (1998); Chandran et al., Indian J Exp Biol. 35(8):801-809 (1997); Margalit, Crit Rev Ther Drug Carrier Syst. 12(2-3):233-261 (1995); U.S. Patent Nos. 5,567,434; 5,552,157; 5,565,213; 5,738.868; and 5,795,587).

Liposomes have been used successfully with a number of cell types that 10 are normally resistant to transfection by other procedures including T cell suspensions, primary hepatocyte cultures and PC 12 cells (Renneisen et al., J Biol Chem. 265(27):16337-16342 (1990); Multer et al., DNA Cell Biol. 9(3):221-229 (1990)). In addition, liposomes are free of the DNA length constraints that are typical of viral-based delivery systems. Linosomes have been used effectively to introduce genes, drugs (ffeath and Martin, Chem Phys Lipids 40(2-4):347-358 (1986); Heath et al., Biochim Biophys Acta. 862(1):72-80 (1986); Balazsovits et al., Cancer Chemother Pharmacol. 23(2):81-6. (1989); Fresta and Puglisi, J. Drug Target 4(2):95-101 (1996)), radiotherapeutic agents (Pikol et al., Arch Surv. 122(12):1417-1420 (1987)), enzymes (Imaignosi et al., Stroke 21(9):1312-1317 (1990); Imaizumi et al., Acta Neurockir Suppl (Wien) 51:236-238 (1990)), viruses (Faller and Baltimore, J Virol, 49(1):269-272 (1984)), transcription 20 factors and allosteric effectors (Nicolan and Gersonde, Naturwissenschaften 66(11):563-566 (1979)) into a variety of cultured cell lines and animals. In addition, several successful clinical trails examining the effectiveness of linosome-mediated drug delivery have been completed (Lopez-Berestein et al., J Infect Dis. 151(4):704-710 (1985); Lopez-25 Berestein et al., Cancer Drue Delly, 2(3):183-189 (1985); Coune, Infection 16(3):141-147 (1988): Sculjer et al., Eur. J. Cancer Clin. Oncol. 24(3):527-38 (1988)). Perthermore, several studies suggest that the use of liposomes is not associated with autoimmune responses, toxicity or gonadal localization after systemic delivery (Mori and Pukatsu, Epilepsia 33(6):994-1000 (1992)).

Liposomes are formed from phospholipids that are dispersed in an aqueous medium and spontaneously form multilamellar concentric bilayer vesicles (also termed multilamellar vesicles (MLVs). MLVs generally have diameters of from 25 nm to 4 nm.

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Someation of MLVs results in the formation of small unilamellar vesicles (SUVs) with diameters in the range of 200 to 500 Å, containing an aqueous solution in the core.

Liposomes bear reaemblance to cellular membranes and are contemplated for use in connection with the present invention as carriers for the peptide compositions. They are widely suitable as both water- and lipid-soluble substances can be entrapped, i.e. in the aqueous spaces and within the bilayer itself, respectively. It is possible that the drug-bearing liposomes may even be employed for site-specific delivery of active agents by selectively modifying the liposomal formulation.

In addition to the teachings of Couvreur et al. (1977), supra; Couvreur et al. (1988), supra), the following information may be utilized in generating liposomal formulations. Phospholipids can form a variety of structures other than liposomes when dispersed in water, depending on the molar ratio of lipid to water. At low ratios the liposome is the preferred structure. The physical characteristics of liposomes depend on pH, ionic strength and the presence of divalent cations. Liposomes can show low permeability to ionic and polar substances, but at elevated temperatures undergo a phase transition which markedly alters their permeability. The phase transition involves a change from a closely packed, ordered structure, known as the gel state, to a loosely packed, less-ordered structure, known as the fluid state. This occurs at a characteristic phase-transition temperature and results in an increase in permeability to ions, sugars and drugs.

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In addition to temperature, exposure to proteins can alter the permeability of liposomes. Certain soluble proteins, such as cytochrome c, bind, deform and penetrate the bilayer, thereby causing changes in permeability. Cholesterol inhibits this penetration of proteins, apparently by packing the phospholipids more tightly. It is contemplated that the most useful liposome formations for antibiotic and inhibitor delivery will contain chalesterol.

The ability to trap solutes varies between different types of liposomes. For example, MLVs are moderately efficient at trapping solutes, but SUVs are extremely inefficient. SUVs offer the advantage of homogeneity and reproducibility in size distribution, however, and a compromise between size and trapping efficiency is offered by large unifamellar vesicles (LUVs). These are prepared by other evaporation and are three to four times more efficient as solute entrapment than MLVs.

In addition to liposome characteristics, an important determinant in entrapping compounds is the physicochemical properties of the compound itself. Polar compounds are trapped in the aqueous spaces and nonpolar compounds bind to the lipid bilayer of the vesicle. Polar compounds are released through permeation or when the bilayer is broken, but nonpolar compounds remain affiliated with the bilayer unless it is disrapted by temperature or exposure to lipoproteins. Both types show maximum efflux rates at the phase transition temperature.

Liposomes interact with cells via four different mechanisms: endocytosis by phagocytic cells of the reticuloendothelial system such as macrophages and neutrophils; adsorption to the cell surface, either by nonspecific weak hydrophobic or electrostatic forces, or by specific interactions with cell-surface components; fusion with the plasma cell membrane by insertion of the lipid bilayer of the liposome into the plasma membrane, with simultaneous release of liposomal contents into the cytoplasm; and by transfer of liposomal lipids to cellular or subcellular membranes, or vice versa, without any association of the liposome contents. It often is difficult to determine which mechanism is operative and more than one may operate at the same time.

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The fate and disposition of intravenously injected liposomes depend on their physical properties, such as size, fluidity, and surface charge. They may persist in tissues for hours or days, depending on their composition, and half lives in the blood range from minutes to several hours. Larger liposomes, such as MLVs and LUVs, are taken up rapidly by phagocytic cells of the reticuloendothelial system, but physiology of the circulatory system restrains the exit of such large species at most sites. They can exit only in places where large openings or porce exist in the capillary endothelium, such as the sinusoids of the liver or spleen. Thus, these organs are the prodominate site of uptake. On the other hand, SUVs show a broader tissue distribution but still are sequestered highly in the liver and spleen. In general, this in vivo behavior limits the potential targeting of liposomes to only those organs and tissues accessible to their large size. These include the blood, liver, spleen, bone marrow, and lymphoid organs.

Targeting is cenerally not a limitation in terms of the present invention.

30 However, should specific targeting be desired, methods are available for this to be accomplished. Antibodies may be used to bind to the liposome surface and to direct the antibody and its drug contents to specific antigenic recentors located on a particular cell-

type surface. Carbohydrate determinants (glycoprotein or glycolipid cell-surface components that play a role in cell-cell recognition, interaction and adhesion) may also be used as recognition sites as they have potential in directing liposomes to particular cell types. Mostly, it is contemplated that intravenous injection of liposomal preparations 5 would be used, but other sources of administration are also conceivable.

Alternatively, the invention provides for pharmaceutically-acceptable nanocapsule formulations of the compositions of the present invention. Nanocapsules can generally entrap compounds in a stable and reproducible way (Henry-Michelland et al. (1987); Onintanar-Guerrero et al., Pharm Res. 15(7):1056-1062 (1998); Douglas et al., Crit. Rev. Ther. Drug Carrier Syss. 3(3):233-261 (1987)). To avoid side effects due to intracellular polymeric overloading, such ultrafine particles (sized around 0.1 µm) should be designed using polymers able to be degraded in vivo. Biodegradable polyalkylcyanoacrylate nanoparticles that meet these requirements are contemplated for use in the present invention. Such particles may be are easily made, as described (Couvreur et al., J. Pharm. Sci. 69(2):199-202 (1980); Couvreur et al., (1988), supra; zur Muhlen et al., Eur. J. Pharm. Biopharm. 45(2):149-155 (1998); Zambaux et al., J. Controlled Release 50(1-3):31-40 (1998); Pinto-Alphandry et al. (1995); and U.S. Patent No. 5,145,684).

R. Vaccines

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In certain preferred embodiments of the present invention, vaccines are 20 provided. The vaccines will generally comprise one or more pharmaceutical compositions, such as those discussed above, in combination with a non-specific immune response enhancer. A non-specific immune response enhancer may be any substance that enhances or potentiates an immane response (antibody anti/or cell-mediated) to an exogenous antigen. Examples of non-specific immune response enhancers include 25 adjuvants, biodegradable microspheres (e.g., polylactic galactide) and liposomes (into which the compound is incorporated; see e.g., Fullerton, U.S. Patent No. 4,235,877). Vaccine preparation is generally described in, for example, Powell and Newman, eds., "Vaccine Design (the subunit and adjuvant approach)," Plemem Press (NY, 1995). Vaccines may be designed to generate antibody immunity and/or cellular immunity such as that arising from CTL or CD4+ T cells.

Pharmaceutical compositions and vaccines within the scope of the present invention may also contain other compounds, which may be biologically active or inactive. For example, one or more immunogenic portions of other *Mycobacterium* antigens may be present, either incorporated into a fusion polypeptide or as a separate compound, within the composition or vaccine. Polypeptides may, but need not, be conjugated to other macromolecules as described, for example, within U.S. Patent Nos. 4,372,945 and 4,474,757. Pharmaceutical compositions and vaccines may generally be used for prophylactic and therapeutic purposes.

Illustrative vaccines may contain DNA encoding one or more of the 10 polypeptides as described above, such that the polypeptide is generated in situ. Such a polymacicotide may comprise DNA, RNA, a modified nucleic acid or a DNA/RNA hybrid. As noted above, the nucleic acid may be present within any of a variety of delivery systems known to those of ordinary skill in the art, including nucleic acid expression systems, bacteria and viral expression systems. Numerous gene delivery 15 techniques are well known in the art, such as those described by Rolland, Crit. Rev. Therap, Drug Carrier Systems 15:143-198 (1998), and references eited therein. Appropriate nucleic acid expression systems contain the necessary DNA sequences for expression in the patient (such as a suitable promoter and terminating signal). Bacterial delivery systems involve the administration of a bacterium (such as Bacillus-Calmette-20 Guerrin) that expresses an immunogenic portion of the notypertide on its cell surface or secretes such an epitope. In a preferred embodiment, the DNA may be introduced using a viral expression system (a.e., vaccinia or other pox virus, retrovirus, or adenovirus). which may involve the use of a non-pathogenic (defective), replication competent virus. Suitable systems are disclosed, for example, in Fisher-Hoch et al., Proc. Natl. Acad. Sci. 25 USA 86:317-321 (1989); Flexner et al., Ann. N.Y. Acad. Sci. 569:86-103 (1989); Flexner et al., Vaccine 8:17-21 (1990); U.S. Patent Nos. 4,603,112; 4,769,330; and 5,017,487; WO 89/01973; U.S. Patent No. 4,777,127; GB 2,290,651; EP 0,345,242; WO 91/02805; Berkner, Biotechniques 6:616-627 (1988); Rosenfeld et al., Science 252:431-434 (1991); Kolls et al., Proc. Natl. Acad. Sci. USA 91:215-219 (1994); Kass-Eisler et al., Proc. Natl. Acad. Sci. USA 90:11498-11502 (1993); Guzman et al., Circulation 88:2838-2848 (1993); and Guzman et al., Cir. Res. 73:1202-1207 (1993). Techniques for incorporating DNA into such expression systems are well known to those of ordinary skill in the art.

The DNA may also be "naked," as described, for example, in Ulmer et al., Science 259:1745-1749 (1993) and reviewed by Cohen, Science 259:1691-1692 (1993). The uptake of naked DNA may be increased by coating the DNA onto biodegradable beads, which are efficiently transported into the cells. It will be apparent that a vaccine may comprise both a polymedeotide and a polypeptide component. Such vaccines may provide for an enhanced immune response.

In a related aspect, a DNA vaccine as described supra may be administered simultaneously with or sequentially to either a polypeptide of the present invention or a known Mycobacterium antigen, such as the 38 kD antigen described above For example, administration of DNA encoding a polypeptide of the present invention, either "taked" or in a delivery system as described supra, may be followed by administration of an antigen in order to enhance the protective immune effect of the vaccine.

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It will be apparent that a vaccine may contain pharmaceutically acceptable salts of the polynucleotides and polypeptides provided herein. Such salts may be prepared from pharmaceutically acceptable non-toxic bases, including organic bases (e.g., salts of primary, secondary and tertiary amines and basic amino acids) and inorganic bases (e.g., sodium, potassium, lithium, ammonium, calcium and magnesium salts).

While any suitable carrier known to those of ordinary skill in the art may be employed in the vaccine compositions of this invention, the type of carrier will vary depending on the mode of administration. Compositions of the present invention may be formulated for any appropriate manner of administration, including for example, topical, oral, nesal, intravenous, intracranial, intraperitoneal, subcutaneous or intramuscular administration. For parenteral administration, such as subcutaneous or intramuscular administration, any of the above carriers or a solid carrier, such as mannitol, factose, starch, magnesium carbonate, sodium saccharine, talcum, cellulose, glucose, sucrose, and magnesium carbonate, may be employed. Biodegradable microspheres (e.g., polylactate polyglycolate) may also be employed us carriers for the pharmaceutical compositions of this invention. Suitable biodegradable microspheres are disclosed, for example, in U.S. Patent Nos. 4,897,268; 5,075,109; 5,928,647; 5,811,128; 5,820,883; 5,853,763; 5,814,344 and 5,942,252. One may also employ a carrier comprising the particulate-protein

complexes described in U.S. Patent No. 5,928,647, which are capable of inducing a class I-restricted cytotoxic T lymphocyte responses in a host.

Such compositions may also comprise buffers (e.g., neutral buffered saline or phosphate buffered saline), carbohydrates (e.g., glucose, mannose, sucrose or 5 dextrans), mannitol, proteins, polypeptides or amino acids such as glycine, antioxidants. bacteriostats, chelating agents such as EDTA or glutathione, adjuvants (e.g., aluminum hydroxide), solutes that render the formulation isotonic, hypotonic or weakly hypertunic with the blood of a recipient, suspending agents, thickening agents and/or preservatives. Alternatively, compositions of the present invention may be formulated as a lyophilizate. Compounds may also be encapsulated within liposomes using well known technology.

Any of a variety of immunostimulants may be employed in the vaccines of this invention. For example, an adjuvant may be included. Most adjuvants contain a substance designed to protect the antigen from rapid catabolism, such as aluminum bydroxide or mineral oil, and a stimulator of immune responses, such as lipid A.

Bortadella pertussis at Mycobacterium species at Mycobacterium derived proteins. For

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example, delipidated, deglycolipidated M. vaccae ("pVac") can be used. In another embodiment, BCG is used as an adjuvant. In addition, the vaccine can be administered to a subject previously exposed to BCG. Suitable adjuvants are commercially available as. for example, Freund's Incomplete Adjuvant and Complete Adjuvant (Difco Laboratories, Detroit, MI): Merck Adiuvant 65 (Merck and Company, Inc., Rahway, NJ): AS-2 and 20 derivatives thereof (SmithKline Beecham, Philadelphia, PA); CWS, TDM, Leif, alumianan salts sach as aluminum hydroxide gel (alum) or aluminum phosphate: salts of calcium, iron or zinc; an insoluble suspension of acylated tyrosine; acylated sugars; cationically or anionically derivatized polysaccharides; polyphosphazenes; biodegradable microsoberes; monophosphoryl lipid A and quit A. Cytokines, such as GM-CSF or 25 interleukin-2, -7, or -12, may also be used as adjuvants.

Within the vaccines provided herein, the adjuvant composition is preferably designed to induce an immane response predominantly of the Th1 type. High levels of Th1-type cytokines (e.g., IFN-y, TNFo., IL-2 and IL-12) tend to favor the 30 induction of cell mediated immune responses to an administered antigen. In contrast, high levels of Th2-type cytokines (e.g., IL-4, IL-5, IL-6 and IL-10) tend to favor the induction of humoral immune responses. Following application of a vaccine as provided

herein, a patient will support an immune response that includes Th1- and Th2-type responses. Within a preferred embodiment, in which a response is predominantly Th1-type, the level of Th1-type cytokines will increase to a greater extent than the level of Th2-type cytokines. The levels of these cytokines may be readily assessed using standard assays. For a review of the families of cytokines, see Mosmann & Coffman, Ann. Rev. Immunol. 7:145-173 (1989).

Preferred adjavants for use in eliciting a predominantly Th1-type response include, for example, a combination of monophosphoryl lipid A, preferably 3-de-O-acylated monophosphoryl lipid A (3D-MPL), together with an aluminum salt. MPL adjavants are available from Corixa Corporation (Seattle, WA; see US Patent Nos. 4,436,727; 4,877,611; 4,866,034 and 4,912,094). CpG-containing ofigomucleotides (in which the CpG dinucleotide is unmethylated) also induce a predominantly Th1 response. Such oligonucleotides are well known and are described, for example, in WO 96/02555, WO 99/33488 and U.S. Patent Nos. 6,008,200 and 5,856,462. Immunostimulatory DNA sequences are also described, for example, by Sato et al., Science 273:352 (1996). Another preferred adjavant comprises a saponin, such as Quil A, or derivatives thereof, including QS21 and QS7 (Aquila Biopharmaceuticals Inc., Framingham, MA); Escin; Digitonin; or Gypsophila or Chenopodium quinoa saponius. Other preferred formulations include more than one seponin in the adjavant combinations of the present invention, for example combinations of at least two of the following group comprising QS21, QS7, Quil A, 8-escin, or digitonin, or digitonin.

Alternatively the saponin formulations may be combined with vaccine vehicles composed of chitosan or other polycationic polymers, polylactide and polylactide-co-glycolide particles, poly-N-acetyl glucosamine-based polymer matrix, particles composed of polysaccharides or chemically modified polysaccharides, liposomes and lipid-based particles, particles composed of glycerol monoesters, etc. The saponins may also be formulated in the presence of cholesterol to form particulate structures such as liposomes or ISCOMs. Furthermore, the saponins may be formulated together with a polyoxycthylene ether or ester, in either a non-particulate solution or suspension, or in a particulate structure such as a paucilametar liposome or ISCOM. The saponins may also be formulated with excipients such as Carbopol[®] to increase viscosity, or may be formulated in a dry powder form with a powder excipient such as lactose.

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In one preferred embodiment, the adjuvant system includes the combination of a monophosphoryl lipid A and a saponin derivative, such as the combination of QS21 and 3D-MPL® adjuvent, as described in WO 94/00153, or a less reactogenic composition where the OS21 is quenched with cholesterol, as described in WO 96/33739. Other preferred formulations comprise an oil-in-water emulsion and tocopherol. Another particularly preferred adjavant formulation employing OS21, 3D-MPL® adiavant and tocopherol in an oil-in-water emulsion is described in WO 95/17210.

Another enhanced adjuvant system involves the combination of a CpGcontaining oligonucleotide and a saponin derivative particularly the combination of CpG and QS21 as disclosed in WO 00/09159. Preferably the formulation additionally comprises an oil in water ensulsion and tocopherol.

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Other preferred adjuvants include Montanide ISA 720 (Seppic, France), SAF (Chiron, California, United States), ISCOMS (CSL), MF-59 (Chiron), the SBAS series of adjuvants (e.g., SBAS-2, AS2', AS2," SBAS-4, or SBAS6, available from SmithKline Beecham, Rixensart, Belgium), Detux (Corixa, Hamilton, MT), RC-529 15 (Corixa, Hamilton, MT) and other aminoalkyl glucosaminide 4-phosphates (AGPs), such as those described in pending U.S. Patent Application Serial Nos. 08/853,826 and 09/074,720, the disclosures of which are incorporated berein by reference in their entireties, and polyoxyethylene ether adjavants such as those described in WO 99/52549A1 20

Other preferred adjuvants include adjuvant molecules of the general formula (f): HO(CH-CH-O),-A-R. wherein, n is 1-50. A is a bond or -C(O)-, R is C1.st affect or Phenyl C1.st affect.

One embodiment of the present invention consists of a vaccine formulation 25 comprising a polyoxyethylene other of general formula (f), wherein n is between 1 and preferably 4-24, most preferably 9; the R component is C_{1.00}, preferably C₂-C₂₀ alkyl and most preferably C_{12} alkyl, and A is a bond. The concentration of the polyoxyethylene ethers should be in the range 0.1-20%, preferably from 0.1-10%, and most preferably in the range 0.1-1%. Preferred polyoxyethylene others are selected from the following group: polyoxyethylene-9-lauryl other, polyoxyethylene-9-steoryl ether, polyoxyethylene-8-steoryl ether, polyoxyethylene-4-huryl ether, polyoxyethylene-35-lanzyl ether, and polyoxyethylene-23-lauryl ether. Polyoxyethylene ethers such as polyoxyethylene lauryl

ether are described in the Merck index (12th edition; entry 7717). These adjuvant molecules are described in WO 99/52549.

The polyoxyethylene other according to the general formula (I) above may, if desired, be combined with another adjuvant. For example, a preferred adjuvant combination is preferably with CpG as described in the pending UK patent application GB 9820956.2

Any vaccine provided herein may be prepared using well known methods that result in a combination of antigen, immune response cahancer and a suitable carrier or excipient. The compositions described herein may be administered as part of a sustained release formulation (i.e., a formulation such as a capsule, sponge or gel (composed of polysaccharides, for example) that effects a slow release of compound following administration). Such formulations may generally be prepared using well known technology (see, e.g., Coombes et al., Vaccine 14:1429-1438 (1996)) and administered by, for example, oral, rectal or subcutaneous implantation, or by 15 implantation at the desired target site. Sustained-release formulations may contain a polypeptide, polynucleotide or antibody dispersed in a carrier matrix and/or contained within a reservoir surrounded by a rate controlling membrane.

Carriers for use within such formulations are biocompatible, and may also be biodegradable; preferably the formulation provides a relatively constant level of active component release. Such carriers include microparticles of poly(lactide-co-glycolide). polyacrylate, latex, starch, cellulose, dextran and the like. Other delayed-release carriers include supramolecular biovectors, which comprise a non-liquid hydrophilic core (e.g., a cross-linked polysaccharide or oligosaccharide) and, optionally, an external layer comprising an amphiphilic compound, such as a phospholipid (see e.g., U.S. Patent No. 25 5,151,254 and PCT applications WO 94/20078, WO/94/23701 and WO 96/06638). The amount of active compound contained within a sustained release formulation depends upon the site of implantation, the rate and expected duration of release and the nature of the condition to be treated or prevented.

C. Delivery vehicles

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Any of a variety of delivery vehicles may be employed within pharmaceutical compositions and vaccines to facilitate production of an antigen-specific immune response that targets infected cells. Delivery vehicles include antigen presenting

cells (APCs), such as dendritic cells, macrophages, B cells, monocytes and other cells that may be engineered to be efficient APCs. Such cells may, but need not, be genetically modified, e.g., to increase the capacity for presenting the antigen, to improve activation and/or maintenance of the T cell response and/or to be immunologically compatible with the receiver (i.e., matched HLA haplotype). APCs may generally be isolated from any of a variety of biological fluids and organs and may be autologous, allogeneic syngeneic or xenogeneic cells.

Certain preferred embodiments of the present invention use dendritic cells or progenitors thereof as antigen-presenting cells. Dendritic cells are highly potent APCs (Banchereau and Steinman, Nature 392:245-251 (1998)) and have been shown to be effective as a physiological adjuvant for eliciting prophylactic or therapeutic immunity (see Timmerman and Levy, Ann. Rev. Med. 50:507-529 (1999)). In general, dendritic cells may be identified based on their typical shape (stellate in situ, with marked cytoplasmic processes (dendrites) visible in vibra), their ability to take up process and present antigeus with high efficiency and their ability to activate naïve T cell responses. 15 Dendritic cells may, of course, be engineered to express specific cell-surface recentors or ligands that are not commonly found on dendritic cells in vivo or ex vivo, and such modified dendritic cells are contemplated by the present invention. As an alternative to dendritic cells, secreted vesicles antigen-loaded dendritic cells (called exosomes) may be used within a vaccine (see Zitvogel et al., Nature Med. 4:594-600 (1998)).

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Dendritic cells and progenitors may be obtained from peripheral blood, bone marrow, lymph nodes, spleen, skin, umbilical cord blood or any other suitable tissue or fluid. For example, dendritic cells may be differentiated as vivo by adding a combination of cytokines such as GM-CSF, IL-4, IL-13 and/or TNFo to cultures of monocytes harvested from peripheral blood. Alternatively, CD34 positive cells harvested from peripheral blood, umbilical cord blood or bone marrow may be differentiated into dendritic cells by adding to the culture medium combinations of GM-CSF, IL-3, TNF a. CD40 ligand, LPS, flt3 ligand and/or other compound(s) that induce materation and proliferation of dendritic cells.

Dendritic cells are conveniently categorized as "immature" and "mature" cells, which allows a simple way to discriminate between two well characterized phonotypes. However, this nomenclature should not be construed to exclude all possible

intermediate stages of differentiation. Immature dendritic cells are characterized as APC with a high capacity for satigen uptake and processing, which correlates with the high expression of Fey recenter and mannose receptor. The mature phenotype is typically characterized by a lower expression of these markers, but a high expression of cell surface molecules responsible for T cell activation such as class I and class II MHC, adhesion molecules (e.g., CD54 and CD11) and costimulatory molecules (e.g., CD40, CD80, CD86 and 4-1BB).

APCs may generally be transfected with a polynucleotide encoding a Mycobacterium antigen (or portion or other variant thereof) such that the Mycobacterium 10 polypeptide, or an immunogenic portion thereof, is expressed on the cell surface. Such transfection may take place ex vivo, and a composition or vaccine comprising such transfected cells may then be used for therapeutic purposes, as described herein. Alternatively, a gene delivery vehicle that targets a dendritic or other antigen presenting cell may be administered to a patient, resulting in transfection that occurs in vivo. In vivo and ax vivo transfection of dendritic cells, for example, may generally be performed using say methods known in the art, such as those described in g.g., WO 97/24447, or the gene gun approach described by Mahvi et al., Immunology and cell Biology 75:456-460 (1997). Astrigen loading of deadritic cells may be achieved by incubating deadritic cells or progenitor cells with the Mycobacterium polypeptide, DNA (naked or within a plasmid vector) or RNA; or with antigen-expressing recombinant bacterium or viruses (e.g., vaccinia, fowlpox, adenovirus or lentivirus vectors). Prior to londing, the polyneptide may be covalently conjugated to an immanological partner that provides T cell help (e.g., a carrier molecule). Alternatively, a dendritic cell may be pulsed with a non-conjugated immanological partner, separately or in the presence of the polypertide.

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Đ. Therapeutic applications of the compositions of the invention

In further aspects of the present invention, the compositions described supra may be used for immunotherapy of Mycobacterium infection, and in particular tuberculosis. Within such methods, pharmaceutical compositions and vaccines are typically administered to a patient to either prevent the development of Mycobacterium infection or to treat a patient afflicted with Mycobacterium infection. Mycobacterium infection may be diagnosed using criteria generally accepted in the art, such as, e.g., in

the case of tuberculosis, fever, acute inflammation of the lung and/or non-productive cough. Pharmaceutical compositions and vaccines may be administered either prior to or following a treatment such as administration of conventional drags. Administration may be by any suitable route, including, e.g., intravenous, intraperitoneal, intramuscular, subcutaneous, intranasal, intradermal, oral, etc.

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Within certain embodiments, immunotherapy may be active immunotherapy, in which treatment relies on the in vivo stimulation of the endogenous host immune system to react against Mycobacterium infection with the administration of immune response-modifying agents (such as polypeptides and polypacleotides as provided herein).

Within other embodiments, immunotherapy may be passive immunotherapy, in which treatment involves the delivery of agents with established Mycobacterium-immune reactivity (such as effector cells or antibodies) that can directly or indirectly mediate anti-Mycobacterium infection effects and do not necessarily depend on an intact host immune system. Examples of effector cells include T cells as discussed above, T lymphocytes (such as CD8" cytotoxic T lymphocytes and CD4" T-helper tumorinfiltrating lymphocytes), killer cells (such as Natural Killer cells and lymphokineactivated killer cells), B cells and antigen-presenting cells (such as dendritic cells and macrophages) expressing a polypeptide of the invention. T cell receptors and antibody receptors specific for the polypeptides recited herein may be cloned, expressed and transferred into other vectors or effector cells for adoptive immunotherapy. The polypeptides provided herein may also be used to generate antibodies or anti-idiotypic antibodies (as described above and in U.S. Patent No. 4,918,164) for passive immunotherapy.

Effector cells may generally be obtained in sufficient quantities for adoptive immunotherapy by growth in vitro, as described herein. Culture conditions for expanding single antigen-specific effector colls to several billion in number with retention of antigen recognition in vivo are well known in the art. Such in vitro culture conditions typically use intermittent stimulation with autigon, often in the presence of cytokines 30 (such as IL-2) and non-dividing feeder cells. As noted above, immunoreactive polypeptides as provided herein may be used to rapidly expand antigen-specific T cell cultures in order to generate a sufficient number of cells for immunotherapy. In

particular, antigen-presenting cells, such as dendritic, macrophage or B cells, may be pulsed with immunoreactive polypeptides or transfected with one or more polymucleotides using standard techniques well known in the art. For example, antigen-presenting cells can be transfected with a polymucleotide having a promoter appropriate for increasing expression in a recombinant virus or other expression system. Cultured effector cells for use in therapy must be able to grow and distribute widely, and to survive long term in vivo. Studies have shown that cultured effector cells can be induced to grow in vivo and to survive long term in substantial numbers by repeated stimulation with antigen supplemented with IL-2 (see, for example, Cheever et al., Immunological Reviews 157:177, (1997)).

Alternatively, a vector expressing a polypeptide recited herein may be introduced into antigen presenting cells taken from a patient and clonally propagated ex vivo for transplant back into the same patient. Transfected cells may be reintroduced into the patient using any means known in the art, preferably in sterile form by, e.g., injection, intranasal or oral administration.

E. Formulation and administration

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Vaccines and pharmaceutical compositions may be presented in unit-dose or multi-dose containers, such as sealed ampoules or vials. Such containers are preferably hermetically sealed to preserve sterility of the formulation until use. In general, formulations may be stored as suspensions, solutions or emulsions in oily or aqueous vehicles. Alternatively, a vaccine or pharmaceutical composition may be stored in a freeze-dried condition requiring only the addition of a sterile liquid carrier immediately prior to use.

Routes and frequency of administration, as well as dosage, may vary from individual to individual and may parallel those currently being employed in immunization using BCG. In general, the pharmacentical compositions and vaccines may be administered, e.g., by injection (e.g., intracutaneous, intramuacular, intravenous or subcutaneous), intramasally (e.g., by aspiration) or orally. Between 1 and 3 doses may be administered for a 1-36 week period. Preferably, 3 doses are administered, at intervals of 3-4 months, and booster vaccinations may be given periodically thereafter. Alternate protocols may be appropriate for individual patients. A suitable dose is an amount of polypeptide or DNA that, when administered as described supra, is capable of raising an

immune response in an immunized patient sufficient to protect the patient from
Mycobacterium infection for at least 1-2 years. When used for a therapeutic purpose, a
suitable dose is the amount that is capable of raising and immune response in a patient
that is sufficient to obtain an improved clinical outcome (e.g., more frequent cure) in
treated patients as compared to non-treated patients. Increases in preexisting immune
responses to a Mycobacterium protein generally correlate with an improved clinical
outcome. Such immune responses may generally be evaluated using standard
proliferation, cytotoxicity or cytokine assays, which may be performed using samples
obtained from a patient before and after treatment.

In general, the amount of polypeptide present in a dose (or produced in situby the DNA in a dose) ranges from about 1 pg to about 100 mg per kg of host, typically from about 10 pg to about 1 mg, and preferably from about 100 pg to about 1 µg. Suitable dose sizes will vary with the size of the patient, but will typically range from about 0.1 ml to about 5 ml.

F. Diagnostic kits

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The present invention further provides kits for use within any of the above diagnostic methods. Such kits typically comprise two or more components uccessary for performing a diagnostic assay. Components may be compounds, reagents, containers and/or equipment. For example, one container within a kit may contain a monoclonal antibody or fragment thereof that specifically binds to a Mycobacterium antigen. Such autibodies or fragments may be provided attached to a support material, as described above. One or more additional containers may enclose elements, such as reagents or buffers, to be used in the assay. Such kits may also, or alternatively, contain a detection reagent as described above that contains a reporter group suitable for direct or indirect detection of antibody binding.

Alternatively, a kit may be designed to detect the level of mRNA encoding a Mycobacterium antigen in a biological sample. Such kits generally comprise at least one oligonucleotide probe or primer, as described above, that hybridizes to a polynucleotide encoding a Mycobacterium antigen. Such an oligonucleotide may be used, for example, within a PCR or hybridization assay. Additional components that may be present within such kits include a second oligonucleotide and/or a diagnostic reagent

or container to facilitate the detection of a polynucleotide encoding a Mycobacterium antisen.

All publications and patent applications cited in this specification are

herein incorporated by reference as if each individual publication or patent application
were specifically and individually indicated to be incorporated by reference.

Although the foregoing invention has been described in some detail by way of illustration and example for purposes of clarity of understanding, it will be readily apparent to one of ordinary skill in the art in light of the teachings of this invention that certain changes and modifications may be made thereto without departing from the spirit or scope of the appended claims.

VIII. EXAMPLES

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EXAMPLE

PURIFICATION AND CHARACTERIZATION OF M. TUBERCULOSIS POLYPEPTIDES USING CD4+ T CELL LINES GENERATED FROM HUMAN PBMC

M. tuberculosis antigens of the present invention were isolated by expression cloning of eDNA libraries of M. tuberculosis strains H37Rv and Erdman easentially as described by Sanderson et al. (J. Exp. Med., 182:1751-1757 (1995)) and were shown to induce PBMC proliferation and IFN-y in an immunoreactive T cell line.

Two CD4+ T cell lines, referred to as DC-4 and DC-5, were generated against dendritic cells infected with M. tuberculosis. Specifically, dendritic cells were prepared from adherent PBMC from a single donor and subsequently infected with tuberculosis. Lymphocytes from the same donor were cultured under limiting dilution conditions with the infected dendritic cells to generate the CD4+ T cell lines DC-4 and DC-5. These cell lines were shown to react with crude soluble proteins from M. tuberculosis but not with Tb38-1. Limiting dilution conditions were employed to obtain a third CD4+ T cell line, referred to as DC-6, which was shown to react with both crude soluble proteins and Tb38-1.

Genomic DNA was isolated from the M. tuberculosis strains H37Rv and Erdman and used to construct expression libraries in the vector pBSK(-) using the

Lambda ZAP expression system (Stratagene, La Jolla, CA). These libraries were transformed into E. coli, pools of induced E. coli cultures were incubated with dendritic cells, and the ability of the resulting incubated dendritic cells to stimulate cell proliferation and IFN-7 production in the CD4+ T cell line DC-6 was examined as described below in Example 2. Positive pools were fractionated and re-tested until pure M. tubercularis planes were obtained.

Nineteen clones were isolated, of which nine were found to contain the previously identified M. tuberculosis antigens TbH-9 and Tb38-1, disclosed in U.S. Patent Application No. 08/533,634. The determined cDNA sequences for the remaining ten clones (hereinafter referred to as Tb224, Tb636, Tb424, Tb436, Tb398, Tb508, Tb441, Tb475, Tb488 and Tb465) are provided in SEQ ID NO:1-10, respectively. The corresponding predicted amino acid sequences for Tb224 and Tb636 are provided in SEQ ID NO:13 and 14, respectively. The open reading frames for these two antigens were found to show some homology to TbH-9. Tb224 and Tb636 were also found to be overlapping clones.

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Tb424, Tb436, Tb398. Tb508, Tb441, Tb475, Tb488 and Tb465 were each found to contain two small open reading frames (referred to as ORF-1 and ORF-2) or truncated forms thereof, with minor variations in ORF-1 and ORF-2 being found for each clone. The predicted amino acid sequences of ORF-1 and ORF-2 for Tb424, Tb436, Tb598, Tb508, Tb441, Tb475, Tb488 and Tb465 are provided in SEQ ID NO:16 and 17, 18 and 19, 20 and 21, 22 and 23, 24 and 25, 26 and 27, 28 and 29, and 30 and 31, respectively. In addition, clones Tb424 and Tb436 were found to contain a third apparent open reading frame, referred to as ORF-U. The predicted amino acid sequences of ORF-U for Tb424 and Tb436 are provided in SEQ ID NO:32 and 33, respectively. Tb424 and Tb436 were found to be either overlapping clones or recently duplicated/transposed copies. Similarly Tb398, Tb508 and Tb465 were found to be either overlapping clones or recently duplicated/transposed copies. as were Tb475 and Tb488.

These sequences were compared with known sequences in publicly
available sequence databases using the BLASTN system. No homologies to the antigens
Tb224 and Tb431 were found. Tb636 was found to be 100% identical to a cosmid
previously identified in M. tuberculosis. Similarly, Tb508, Tb488, Tb398, Tb424, Tb436,
Tb441, Tb465 and Tb475 were found to show homology to known M. tuberculosis

cosmids. In addition, Tb488 was found to have 100% boundary to M. tuberculosis topolsomerase I.

Seventeen overlapping peptides to the open reading frames ORF-1 (referred to as 1-1 - 1-17; SEQ ID NO:34-50, respectively) and thirty overlapping peptides to the open reading frame ORF-2 (referred to as 2-1 - 2-30, SEQ ID NO:51-80, respectively) were synthesized using the procedure described below in Example 4.

The ability of the synthetic peptides and of recombinant ORF-1 and ORF-2 to induce T cell proliferation and IFN-γ production in PBMC from PPD-positive donors was assayed as described below in Example 2. Figs. 1A-B and 2A-B illustrate stimulation of T cell proliferation and IFN-γ by recombinant ORF-2 and the synthetic peptides 2-1 - 2-16 for two donors, referred to as D7 and D160, respectively.

Recombinant ORF-2 (referred to as MTI) stimulated T cell proliferation and IFN-γ production in PBMC from both donors. The amount of PBMC stimulation seen with the individual synthetic peptides varied with each donor, indicating that each donor recognizes different epitopes on ORF-2. The proteins encoded by ORF-1, ORF-2 and ORF-U were subsequently named MTS. MTI and MSF, respectively.

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Eighteen overlapping peptides to the sequence of MSF (referred to as MSF-4-MSF-48; SEQ ID NO:84-101, respectively) were synthesized and their ability to stimulate T cell proliferation and IFN-y production in a CD4+ T cell line generated against M. tuberculosis culture filtrate was examined as described below. The peptides referred to as MSF-12 and MSF-13 (SEQ ID NO:95 and 96, respectively) were found to show the highest levels of reactivity.

Two overlapping peptides (SEQ ID NO:81 and 82) to the open reading frame of Tb224 were synthesized and shown to induce T cell proliferation and IFN-γ production in PBMC from PPD-positive donors.

Two CD4+ T cell lines from different donors were generated against M. tuberculoxis infected dendritic cells using the above methodology. Screening of the M. tuberculosis cDNA expression library described above using this cell line, resulted in the isolation of two clones referred to as Tb867 and Tb391. The determined cDNA sequence for Tb867 (SEQ ID NO:102) was found to be identical to the previously isolated M. tuberculosis cosmid SCY22Gi0, with the candidate resetive open reading frame encoding

a 750 amino acid M. tuberculosis protein kinase. Comparison of the determined cDNA sequence for Tb391 (SEQ ID NO:103) with those in publicly available sequence databases revealed no significant homologies to known sequences.

In further studies, CD4+ T cell lines were generated against M.

5. tuberculosis culture filtrate, essentially as outlined above, and used to screen the M.

tuberculosis Frdman cDNA expression library described above. Five reactive clones, referred to as Tb431, Tb472, Tb470, Tb638 and Tb962 were isolated. The determined cDNA sequences for Tb431, Tb472, Tb470, and Tb838 are provided in SEQ ID NO:11, 12, 104 and 105, respectively, with the determined cDNA sequences for Tb962 being provided in SEQ ID NO:106 and 107. The corresponding predicted amino acid sequence for Tb431 is provided in SEQ ID NO:15.

Subsequent studies led to the isolation of a full-length cDNA sequence for Tb472 (SEO ID NO:168). Overlapping pentides were synthesized and used to identify the reactive open reading frame. The predicted amino acid sequence for the protein 15 encoded by Tb472 (referred to as MSL) is provided in SEO ID NO:109. Comparison of the sequences for Tb472 and MSL with those in publicly available sequence databases as described above, revealed no homologies to known sequences. Fifteen overlapping peptides to the sequence of MSL (referred to as MSL-1 · MSL-15; SEO ID NO:110-124. respectively) were synthesized and their ability to stimulate T cell proliferation and IFN-y 20 production in a CD4+ T cell line generated against M. tuberculosis culture filtrate was examined as described below. The pentides referred to as MSL-10 (SEO ID NO:119) and MSL-11 (SEO ID NO:120) were found to show the highest level of reactivity. Comparison of the determined cDNA sequence for Tb838 with those in publicly available sequence databases revealed identity to the previously isolated M. tuberculoris costaid 25 SCY07H7. Comparison of the determined cDNA sequences for the clone Tb962 with those in publicly available sequence databases revealed some homology to two previously identified M. tuberculosis cosmids, one encoding a portion of bactoferritin. However, recombinant bactoferritin was not found to be reactive with the T cell line used to implete Tb962.

30 The clone Tb470, described above, was used to recover a full-length open reading frame (SEQ ID NO:125) that showed homology with TbH9 and was found to encode a 40 kDa antigen, referred to as Mth40. The determined amino acid sequence for

Mib40 is provided in SEQ ID NO:126. Similarly, subsequent studies led to the isolation of the full-length cDNA sequence for Tb431, provided in SEQ ID NO:83, which was also determined to contain an open reading frame encoding Mtb40. Tb470 and Tb431 were also found to contain a potential open reading frame encoding a U-ORF-like antigen.

Screening of an M. tuberculosis Erdman cDNA expression library with multiple CD4+ T cell lines generated against M. tuberculosis culture filtrate, resulted in the isolation of three clones, referred to as Tb366, Tb433 and Tb439. The determined cDNA sequences for Tb366, Tb433 and Tb439 are provided in SEQ ID NO:127, 128 and 129, respectively. Comparison of these sequences with those in publicly available sequence databases revealed no significant homologies to Tb366. Tb433 was found to show some homology to the previously identified M. tuberculosis antigen MPTS3. Tb439 was found to show 100% identity to the previously isolated M. tuberculosis cosmid SCY02B10.

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A CD4+ T cell line was generated against M. tuberculosis PPD, essentially described above, and used to screen the above M. tuberculosis Erdman cDNA expression library. One reactive clone (referred to as Tb372) was isolated, with the determined cDNA sequences being provided in SEQ ID NO:130 and 131. Comparison of these sequences with those in publicly available sequence databases revealed no significant homologies.

In further studies, screening of an *M. tuberculosis* cDNA expression library with a CD4+ T cell line generated against dendritic cells that had been infected with suberculosis for 8 days, as described above, led to the isolation of two clones referred to as Th390R5C6 and Th390R2C11. The determined cDNA sequence for Tb390R5C6 is provided in SEQ ID NO:132, with the determined cDNA sequences for Th390R2C11 being provided in SEQ ID NO:133 and 134. Th390R5C6 was found to show 100% identify to a previously identified *M. nuberculosis* cosmid.

In subsequent studies, the methodology described above was used to screen an M. tuberculosis genomic DNA library prepared as follows. Genomic DNA from M. tuberculosis Erdman strain was randomly sheared to an average size of 2 kb, and blunt ended with Klenow polymerase, followed by the addition of EcoRI adaptors. The insert was subsequently ligated into the Screen phage vector (Novagen, Madison, WI) and packaged in vitro using the PhageMaker extract (Novagen). The phage library (referred to

as the Erd \(\) Screen library) was amplified and a portion was converted into a plasmid expression library by an autosubcloning mechanism using the \(E. \) coli strain BM25.8 (Novagen). Plasmid DNA was purified from BM25.8 cultures containing the pSCREEN recombinants and used to transform competent cells of the expressing host strain BL2t(DE3)pLysS. Transformed cells were aliquoted into 96 well microtiter plates with each well containing a pool size of approximately 50 colonies. Replica plates of the 96 well plasmid library format were induced with IPTG to allow recombinant protein expression. Following induction, the plates were centrifuged to pellet the \(E. \) coli which was used directly in T cell expression cloning of a CD4+T cell line prepared from a PPD-positive donor (donor 160) as described above. Pools containing \(E. \) coli expressing \(M. \) tuberculosis T cell antigens were subsequently broken down into individual colonies and reassaved in a similar fashion to identify positive hits.

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Screening of the T cell line from donor 160 with one 96 well plate of the End a Screen library provided a total of nine positive hits. Previous experiments on the screening of the pBSK library described above with T cells from donor 160 suggested 15 that most or all of the positive clones would be TbH-9. Tb38-i or MTI (disclosed in U.S. Patent Application No. 08/533,634) or variants thereof. However, Southern analysis revealed that only three wells hybridized with a mixed probe of TbH-9. Tb38-1 and MTI. Of the remaining six positive wells, two were found to be identical. The determined 5' 20 cDNA segmences for two of the isolated clones (referred to as YI-26C1 and YI-86C11) are provided in SEQ ID NO:135 and 136, respectively. The full length cDNA sequence for the isolated clone referred to as hTcc#l is provided in SEQ ID NO:137, with the corresponding predicted amino acid sequence being provided in SEQ ID NO:138. Comparison of the sequences of hTee#1 to those in publicly available sequence databases 28 as described above, revealed some homology to the previously isolated M. tuberculosis cosmid MTCY07H7B.06.

EXAMPLE 2

INDUCTION OF T CELL PROLIFERATION AND INTERFERON-? PRODUCTION BY M. TUBERCULOSIS ANTIGENS

The ability of recombinant M. tuberculosis antigens to induce T-cell proliferation and interferon-q production may be determined as follows.

Proteins may be induced by IPTG and purified by gel clution, as described in Skeiky et al., J. Exp. Med. 181:1527-1537 (1995). The purified polypeptides are then screened for the ability to induce T-cell proliferation in IPBMC preparations. The PBMCs from denors known to be PPD skin test positive and whose T-cells are known to 5 proliferate in response to PPD are cultured in medium comprising RPMI 1640 supplemented with 10% pooled human serum and 50 µg/ml gentamicin. Purified polypeptides are added in duplicate at concentrations of 0.5 to 10 µg/ml. After six days of culture in 96-well round-bottom plates in a volume of 200 µl, 50 µl of medium is removed from each well for determination of IFN-7 levels, as described below. The plates are then pulsed with 1 µCl/well of tritiated thymidine for a further 18 hours, harvested and tritium uptake determined using a gas scintillation counter. Fractions that result in proliferation in both replicates three fold greater than the proliferation observed in cells cultured in medium alone are considered positive.

IFN-γ is measured using an enzyme-linked immunosorbent assay (ELISA).

ELISA plates are coaled with a mouse monoclonal antibody directed to human IFN-y (PharMingen, San Diego, CA) in PBS for four hours at room temperature. Wells are then blocked with PBS containing 5% (W/V) non-fat dried milk for 1 hour at room temperature. The plates are washed six times in PBS/0.2% TWEEN-20 and samples diluted 1:2 in culture medium in the ELISA plates are incubated overnight at room 20 temperature. The plates are again washed and a polyclonal rabbit anti-human IFN-y serian diluted 1:3000 in PBS/10% normal goat serian is added to each well. The plates are then incubated for two hours at room temperature, washed and horseradish peroxidase-coupled anti-rabbit IgG (Sigma Chemical So., St. Louis, MO) is added at a 1:2000 dilution in PBS/5% non-fat dried milk. After a further two hour incubation at 25 room temperature, the plates are washed and TMB substrate added. The reaction is stopped after 20 min with 1 N sulfuric acid. Optical density is determined at 450 nm using 570 nm as a reference wavelength. Fractions that result in both replicates giving an OD two fold greater than the mean OD from cells cultured in medium alone, plas 3 standard deviations, are considered positive.

EXAMPLE 3

PURIFICATION AND CHARACTERIZATION OF M. TUBERCULOSIS POLYPEPTIDES USING CD4+ T CELL LINES GENERATED FROM A MOUSE M. TUBERCULOSIS MODEL

Infection of C57BL/6 mice with M. nuberculosis results in the development of a progressive disease for approximately 2-3 weeks. The disease progression is then halted as a consequence of the emergence of a strong protective T cell-mediated immune response. This infection model was used to generate T cell lines capable of recognizing protective M. tuberculosis antigens.

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Specifically, spleen cells were obtained from C57BL/6 mice infected with M. tuberculosis for 28 days and used to raise specific anti-M. tuberculosis T cell lines as described above. The resulting CD4+ T cell lines, in conjunction with normal antigen presenting (spleen) cells from C57BL/6 mice were used to screen the M. tuberculosis Erd \(\frac{1}{2} \) Screen library described above. One of the reactive library pools, which was found to be highly stimulatory of the T cells, was selected and the corresponding active clone (referred to as Y288C10) was isolated.

Sequencing of the clone Y2SSC10 revealed that it contains two potential genes, in tandem. The determined cDNA sequences for these two genes (referred to as mTCC#1 and mTCC#2) are provided in SEQ ID NO:139 and 140, respectively, with the corresponding predicted amino acid sequences being provided in SEQ ID NO:141 and 142, respectively. Comparison of these sequences with those in publicly available sequence databases revealed identity to unknown sequences previously found within the M. tuberculosis cosmid MTY21C12. The predicted amino acid sequences of mTCC#1 and mTCC#2 were found to show some homology to previously identified members of the TbH9 protein family, discussed above.

EXAMPLE 4

SYNTHESIS OF SYNTHETIC POLYPEPTIDES

Polypeptides may be synthesized on a Millipore 9050 peptide synthesizer using FMOC chemistry with HIPTU (O-Benzotriazole-N,N,N',N'-tetramethyluronium hexatiluerophosphate) activation. A Gly-Cys-Gly sequence may be attached to the amino terminus of the peptide to provide a method of conjugation or labeling of the peptide. Cleavage of the peptides from the solid support may be carried out using the following

cleavage mixture: trifluoroacetic acid:ethanedithiolathioautsoler.water:phenol (40:1:2:2:3). After cleaving for 2 hours, the peptides may be precipitated in cold methyl-butyl-ether. The peptide pellets may then be dissolved in water containing 0.1% trifluoroacetic acid (TFA) and lyophilized prior to purification by C18 reverse phase FIPLC. A gradient of 0-60% acetonitrile (containing 0.1% TFA) in water (containing 0.1% TFA) may be used to elute the peptides. Following lyophilization of the pure fractions, the peptides may be characterized using electrospray mass spectrometry and by amino acid analysis.

EXAMPLE 5

USE OF REPRESENTATIVE ANTIGENS FOR SERODIAGNOSIS OF TUBERCULOSIS

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The diagnostic properties of representative M. tuberculosis antigens may be determined by examining the reactivity of antigens with sera from tuberculosisinfected patients and from normal donors as described below.

Assays are performed in 96-well plates coated with 200 ng antigen diluted to 50 μl in carbonate coating buffer, pH 9.6. The wells are coated overnight at 4°C (or 2 hours at 37°C). The plate contents are then removed and the wells are blocked for 2 hours with 200 μl of PBS/1% BSA. After the blocking step, the wells are washed five times with PBS/0.1% Tween 20°v. 50 μl sera, diluted 1:100 in PBS/0.1% Tween 20′0.1% BSA, is then added to each well and incubated for 30 minutes at room temperature. The plates are washed again five times with PBS/0.1% Tween 20°n.

The enzyme conjugate (horseradish peroxidase - Protein A, Zymed, San Francisco, CA) is then 1:10,000 in PBS/0.1% Tween20™/0.1% BSA, and 50 µl of the diluted conjugate is added to each well and incubated for 30 minutes at room temperature. Following incubation, the wells are washed five times with PBS/0.1% Tween 20™. 100 µl of tetramethylbenzidine peroxidase (TMB) substrate (Kirkegaard and Perry Laboratories, Gaithersburg, MD) is added, undiluted, and incubated for about 15 minutes. The reaction is stopped with the addition of 100 µl of 1 NH₂SO₄ to each well, and the plates are read at 450 nm.

EXAMPLE 6

MURINE T CELL EXPRESSION CLONING OF AN MTB ANTIGEN ASSOCIATED WITH THE CONTROL OF TB INFECTION

Genomic DNA form M. tuberculosis Extman strain was randomly sheared 5 to an average size of 2 kb, blunt ended with Klenow polymerase and followed by the addition of EcoRI adaptors. The insert was subsequently ligated into the Screen phage vector predigested with EcoRI (Novagen, Madison, WI) and packaged in vitro using the PhageMaker extract (Novagen, Madison, Wi). The phage library (Erd Screen) was amplified and a portion converted into a plasmid expression library (nScreen) by 10 autosubcloning using the E. coli host strain BM25.8 as suggested by the manufacturer (Novagen, Madison, WI). Plasmid DNA was purified from BM25.8 cultures containing pScreen recombinants and used to transform competent cells of the expressing bost strain BL21(DE3)pl.ysS. Transformed cells were aliquoted into 96 well micro titer plates with each well containing a pool size of ~50 colonies. Replica plates of the 96 well plasmid 15 library format were induced with IPTG to allow recombinant protein expression. Following induction, the plates were centrifuged to pellet the E. coli and the bacterial pellet was resuspended in 200 at of 1 X PBS. The general principle is based on the direct recognition by the T cells of the antigens presented by antigen presenting cells that have internalized a library of E. coli-containing expressed recombinant artigens. The M. 20 suberculosis library was initially divided in pools containing approximately 50-100 transformants/ml distributed in 96-well microtiter plates and stored in a replaca plate manner. Adherent spleen cells were fed with the E. coli pools and incubated for processing for 2 h. After washing the adherent cells were exposed to specific T cell lines in the presence of gentamyoin (50 ug/mi) to inhibit the bacterial growth. T cell 25 recognition of pool containing M. tuberculosis antigens was then detected by proliferation (3H thymidine incorporation). Wells that scored positive were then broken down using the same protocol until a single clone was detected. The gene was then sequenced, subcloned, expressed and the recombinant protein evaluated. Nucleotide sequence comparison of the 0.6 kb insert of clone mTTC#3 with the GenBank database revealed 30 that it is comprised of the amino terminal portion of gene MTV014.03c (locus MTV014;

accession # e1248750) of the Mtb H37Rv strain. The full length nucleotide sequence of mTTC#3 (SBQ ID NO:145) is a 1.86 kb fragment commising the entire ORF with a

predicted molecular weight of ~57 kDa (SEQ ID NO:146). Thus, to maintain consistency with our nomenclature, mTTC#3 is referred to hereafter as MTB57. The full length coding portion of mTTC#3 (MTB57) was PCR amplified using the following primer pairs: 5'(5' ~CAA TTA CAT ATG CAT CAC CAT CAC CAT CAC ATG AAT TAT TCG GTG TTG CCG (SEQ ID NO:147)) and 3' (5' ~CAA TTA AAG CTT TTA GGG CTG ACC GAA GAA GCC (SEQ ID NO:148))h3. The full length nucleic acid coding sequence of mTTC#3 and the corresponding predicted amino acid sequence are provided in Figures 3 and 4, respectively.

EXAMPLE 7

IDENTIFICATION OF MYCOBACTERIUM TUBERCULOSIS ANTIGENS EXCRETED IN URINE OF INFECTED MICE

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Antigen were prepared by infecting intravenously C57BL/6 nice with 4.10⁷ colony forming units (CFU) of M. tuberculosis. 14 days later the animals were blod and their urine was collected in microfuge tubes. Sera were obtained at room temperature. Both sera and urine were centrifuged at 10,000 g for 15 minutes followed by filtration in 0.2u sterile membranes.

Antibodies were produced against the antigens by immunizing normal C57BL/6 mice with either the sera or the urine from the M. tuberculosis infected C57BL/6 mice. The adjuvant used was incomplete Freund's adjuvant (IFA).

CS7BL/6 raice. The adjuvant used was incomplete Freund's adjuvant (IFA). Immunization was carried out according to the following protocol: on day 1, mice were injected in the footpad or in the base of the tail with a mix containing 100 µl of either serum or urine and 100 µl of IFA; on day 14, a mix containing 100 µl of either serum or urine and 100 µl of IFA was injected intraperitoneally to the mice; finally on day 28, either 200 µl of serum or 50 µl of urine were injected to the mice intraperitoneally. By using syngencic mice for the antibody production, only antibodies specific for foreign antigens present in the blood circulation or urine of the C57BL/6 mice, i.e., M. unberculosis antigens, are generated. On day 35, 100 µl of blood were collected by eyebleeding the immunized mice. ELISA assays were performed with the obtained sera using a M. tuberculosis crude lysate. The ELISA experiments revealed that all the mice immunized with either sera or urine from infected donors produced anti-M. tuberculosis antibodies in titers varying from 1/40 to 1/320. No anti-M. tuberculosis antibodies were

found in the sera obtained from the mice before the immunizations.

The antiserum made against the proteins excreted in the urine was used to screen a Mtb expression library prepared in the lambda screen phage expression system. Positive clones were purified and their corresponding inserts sequenced. These inserts were named P1, 2, 3, 4, 6, 7, 8, 9, 10, 11 and 12 (SEQ ID NO:149-159).

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EXAMPLE 8

IDENTIFICATION OF MYCOBACTERIUM TUBERCULOSIS ANTIGENS USING CD4+ T CELL EXPRESSION CLONING

Expression screening using a number of T cell lines generated from

10 healthy PPD-positive individuals has been employed to identify M. tuberculosis clones
encoding reactive antigens. Pools of M. tuberculosis recombinant clones (expressed in E.
colf) were fed to dendritic cells. Autologous T cell lines were incubated with the
dendritic cells and proliferation and INF-gamma production was measured. Reactive
pools were fractionated and re-tested until pure M. tuberculosis clones were achieved.

15 This approach allows for direct screening for T cell antigeus. A related approach has
been used to identify Listeria monocytogenes antigens (see J. Exp. Med. 182:1751-1757
(1995).

From the foregoing, it will be appreciated that, although specific

20 embodiments of the invention have been described herein for the purpose of illustration,
various modifications may be made without deviating from the spirit and scope of the
invention

WHAT IS CLAIMED IS:

ş 1. An isolated polypeptide comprising an amino acid sequence of SEQ ID 2 NO:146, 161, or 163, or an animo acid sequence comprising an immunogenic portion of an 3 amino acid sequence of SEQ ID NO:146, 161, or 163. 2 An isolated polyneptide, wherein said polypeptide is encoded by a ž 2 nacleotide sequence selected from the group consisting of SEO ID NO:145, 149, 156, 151, 152, 153, 154, 155, 156, 157, 158, 159, 160, 162, and 164, or an isolated polypeptide 3 comprising an immunogenic portion of a polypeptide encoded by a nucleotide sequence 4 selected from the group consisting of SEO ID NO:145, 149, 150, 151, 152, 153, 154, 155, 5 6 156, 157, 158, 159, 160, 162, and 164. The polypeptide of claim 1 or 2, wherein the polypeptide is fused to a 2 second polypeotide to form a fusion protein. The fusion protein of claim 3, wherein the two polypeptides are 2 heterologous. S. The fusion protein of claim 3, wherein the polypoptides are Mycobacterium tuberculosis polypeptides. 2 F The fusion protein of claim 3, wherein the second polypeptide is a 2 known Mycobacterium antigen. 1 A polynucleotide comprising a nucleotide sequence encoding a fusion protein according to claim 3. 1 A pharmaceutical composition comprising a fusion protein according 2 to claim 3 and a physiologically acceptable carrier.

 An isolated polynucleotide that specifically hybridizes under moderately stringest conditions to a second polynucleotide comprising a muleotide sequence selected from the group consisting of SEQ ID NO:145, 149, 150, 151, 152, 153, 154, 155,

4 156, 157, 158, 159, 160, 162, and 164.

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*		10.	An isolated polymedeotide that specifically hybridizes under highly		
2	stringent conditions to a second polynucleotide comprising a nucleotide seguence selected				
3	from the group consisting of SEQ ID NO:145, 149, 150, 151, 152, 153, 154, 155, 156, 157,				
4	158, 159, 160, 162, and 164.				
9		[1.	An expression vector comprising a polyancleotide according to claim 5		
2	or 10.				
y		12.	A host cell transformed with an expression vector according to claim		
2	11.				
200		13.	The host cell of claim 12, wherein the host cell is selected from the		
2	group consisting of E. coli, yeast, and mammalian cells.				
Section 1		14.	A method for detecting Mycobacterium infection in a biological		
2	sample, the method comprising the steps of:				
3	(a) contacting a biological sample with at least one polypoptide according to				
4	claim 1 or 2; and				
5		(b) detecting in the sample the presence of antibodies that bind to the			
6	polypeptide, thereby detecting $Mycobacterium$ infection in the biological sample.				
Į		15.	The method of claim 14, wherein the polypeptide is bound to a solid		
2	support.				
1		6.	The method of claims 15, wherein the solid support comprises		
2	nitrocellulose, latex or a plastic material.				
See.		7.	The method of claim 14, wherein the biological sample is selected		
2	from the group consisting of whole blood, sputum, serum, plasma, saliva, cerebrospinal fluid				
3	and urine.				
1		18.	The method of claim 17, wherein the biological sample is whole blood		
2	or sening.				

	19. The method of claim 14, wherein the Mycobucterium infection is a				
	Mycobacterium tuberculosis infection.				
	20. A method for detecting Mycobacterium infection in a biological				
,	sample, the method comprising the steps of:				
,	(a) contacting the sample with at least two oligonucleotide primers, wherein at				
	least one of the oligonucleotide primers specifically hybridizes under stringent conditions to a				
5	polynucleotide according to claim 9; and				
\$	(b) detecting in the sample a polynucleotide sequence that is amplified in the				
7	presence of the oligonucleotide primers, thereby detecting Mycobacterium infection.				
	21. The method of claim 20, wherein the biological sample is selected				
1	from the group consisting of whole blood, sputum, serum, plasma, saliva, cerebrospinal fluid				
,	and arine.				
	22. The method of claim 20, wherein the Mycobacterium infection is a				
2	Mycobacterium tuberculosis infection.				
	23. A method for detecting Mycobacterium infection in a biological				
2	sample, the method comprising the steps of:				
,	(a) contacting the sample with one or more polynucleotide probes that				
,	specifically hybridize to a polynucleotide according to claim 9; and				
,	(b) detecting in the sample a DNA sequence that hybridizes to the				
	oligonucleotide probe, thereby detecting Mycobacterium infection.				
	24. The method of claim 23, wherein the biological sample is selected				
	from the group consisting of whole blood, sputum, serum, plasma, saliva, cerebrospinal fluid				
	sad prine,				
	25. The method of claim 23, wherein the Mycobacterium infection is a				
	Mycobacterium tuberculosis infection.				
	26. A method for detecting Mycobacterium infection in a biological				
	sample, the method comprising the steps of:				

3	(a) c	ontacting the biological sample with a binding agent which is capable of			
4	binding to a polypeptide according to claim 1 or 2; and				
5	(b) detecting in the sample a polypeptide that binds to the binding agent,				
6	thereby detecting Mycobacterium infection in the biological sample.				
No.	27.	The method of claim 26, wherein the binding agent is a monoclonal			
2	antibody.				
l	28.	The method of claim 26, wherein the binding agent is a polyclonal			
2	antibody.				
1	29.	The method of claim 26, wherein the Mycobacterium infection is a			
2	Mycobacterium tuberculosis infection.				
¥.	30.	A diagnostic kit comprising:			
2	(a)	one or more polypeptides according to claim 1 or 2; and			
3	(6)	a detection reagent.			
year.	31.	The kit of claim 30, wherein the polypeptide is immobilized on a solid			
2	support.				
I	32.	The kit of claim 30, wherein the detection reagent comprises a seporter			
2	group conjugated to a binding agent.				
1	33.	The kit of claim 32, wherein the binding agent is selected from the			
2	group consisting of anti-immunoglobulins, Protein G, Protein A and lectins.				
1	34.	The kit of claim 32, wherein the reporter group is selected from the			
2	group consisting of radioisotopes, fluorescent groups, haminescent groups, enzymes, biotin				
3	and dye particles.				
1	35.	A diagnostic kit comprising at least two oligomic leotide primers,			
2	wherein at least one of the oligonecleotide primers specifically hybridizes under stringent				
3	conditions to a polymacleotide according to claim 9.				

3000	36	. A diagnostic kit comprising at least one polynocleotide probe, whereis			
2					
3	the polynucleotide probe specifically hybridizes under stringent conditions to a polynucleotide according to claim 9.				
3	porynuciconare ac	conning to casm 9.			
3	37	. An antibody that binds to a polypeptide according to claim 1 or 2.			
1	38	. The antibody of claim 37, wherein the antibody is a monoclonal			
2	antibody.				
ì	39	. A pharmaceutical composition comprising at least one polypoptide			
2	according to clair	n 1 or 2, and a physiologically acceptable carrier.			
W.	40	A pharmaceutical composition comprising a polynucleotide according			
2	to claim 9 and a physiologically acceptable carrier.				
Va.de	41	The pharmaceutical composition of claim 39 or 40, wherein the			
2	pharmaceutical co	supposition is a vaccine and a non-specific immune response enhancer.			
-	42	The vaccine of claim 41, further comprising a non-specific immune			
2	response enhance	.			
1	43	The vaccine of claim 42, wherein the non-specific immuse enhancer is			
2	an adjuvant.				
1	44	The vaccine of claim 43, wherein the adjuvant is selected from the			
2	group consisting of SBAS-2, QS-21, 3D-MPL, GM-CSF, SAF, ISCOMS, MF-59 and RC-				
3	529.				
· www	45	A method for eliciting or cabancing an immune response to			
2	Mycobacterium in a patient, the method comprising the step of administering to a patient a				
3	pharmaceutical composition according to claims 39 or 40 in an amount effective to elicit or				
4	enhance the immune response.				

46. A method for inhibiting the development of a Mycobacterium infection in a patient, the method comprising the step of administering to a patient an effective amount

2

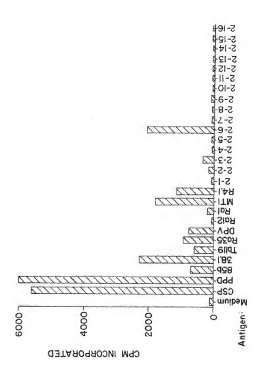
3 of a pharmsceatical composition according to claims 39 or 40, and thereby inhibiting the development of a Mycobacterium infection in the patient. A method for inhibiting the development of a Mycobacterium infection 2 in a patient, the method comprising the step of administering to a patient an effective amount 3 of an antibody according to claim 37, and thereby inhibiting the development of a Mycobacterium infection in the patient. 4 ŝ 48 The method of claims 46 or 47, wherein the Mycobacterium infection is a M. tuberculosis infection. 2 1 49. A method for detecting tuberculosis in a patient, the method 2 comprising the steps of: 3 (a) contacting dermal cells of a patient with at least one polypeptide according to claim 1 or 2; and 4 5 (b) detecting an immune response on the patient's skin and therefrom detecting tuberculosis in the patient. 6 50. The method of claim 49, wherein the immune response is induration. 51. A diagnostic kit comprising: (a) a polypeptide according to claim 1 or 2; and

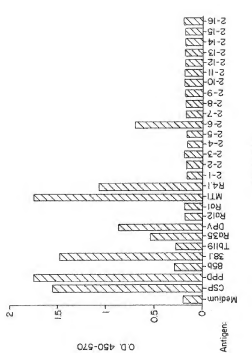
(b)

cells of a patient.

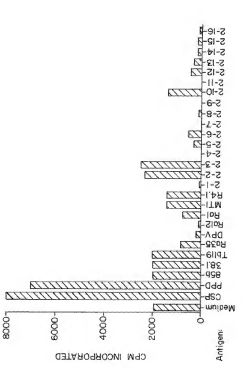
an apparatus sufficient to contact said polypeptide with the demnal





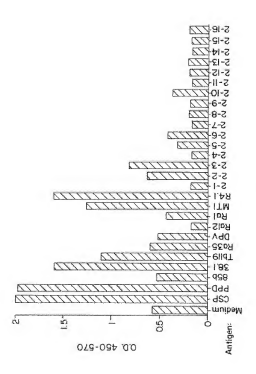


F/G. 18.



SUBSTITUTE SHEET (RULE 26)

F16. 2B.



>mTCCf3His.seg

ATGCATCACCATCACCATCACATGAATTATTCGGTGTTGCCGCCGGAGATTAATTCGTTGCGGATG

GTCTGCCCGATGCTTGCGGCATCGGTGGCTTGGGATGTTTGGCCGCGAGTTGGCGGTGGCGGC GTCCTCGTTTGGGT

GGGTGGTTGGCTGCGGCGGGCGGGCGGGCGGCGGTGGCGCTTGGCTCAGGCCAAGGCGGTGGCCAGT

GCGGGGGGGCGACGGTGCATCCGATGCTGGTGGCGGCCAACCGTAATGCGTTTGTGCAGTTGGTGTT GTCGAATCTGTTTG

GGUAGAATGCGCCGGCGATCGCGGCCGCTGAGGCGATGTATGAACAGATGTGGGCCGCCGATGTGG CCGCGATGCTGGGC

TYGUCAGCTGUGCC
ATCGGCGCGCGCGATCGGCCTCGGCAACATCGGCGGTCGGGAACCTGGGCGGCGGGAACAC

GCGGAAATGACGGT GCCACGAATTTGGGCAGCGGAAATATCGGCAACACCAATCTCGGCAGCGGAAACGTTGGCAATGTC

ARTCTEGGCAGCEG
ARACUGAGGCTTTGGAAACCTCGGCAACGGAAACTTTGGCAGTGGGAACCTCGGCAGTGGAAACAC

CGGAAGTACCAACT

TCGGCGCCGGAAATCTCGGTTCCTTCAACTTGGGCAGTGGAAACATCGCCTCCTACATCGGTT TCGGAAACAACGGC

GRCMATARCCTCGGCCTCGGGAACAATGGCAACAACAACATCGGTTTTGGGCTCACCGGCGACAAC TTGGTGGGCAFTGG

CGCGCTEAACTCGGGCATCGGGAATCTAGGTTTCGGGAACTCGGGTAACAACAACATCGGTTTCTT CAACTCTGGCAACA ACAACGTGGGCTTCTTCAATTCGGGCAACAACAACTTCGGCTTTGGAAACGCGGGGGACATCAACA

ACAACGIGGCTTCTTCAATTCGGGCAACAACGTTCGGCTTTGGAAACGCGGGGGGACATCAACA CGGGCTTCGGAAAC

GCCGGCGACACCACGGGCTTCGGAAACGCCGGCTTCTTCAATATGGGCATCGGGAACGCGGGC AACGAAGACATGG

TAAACGTGGGTTCGCAAACGCGGGCAGTATCAATACGGGATTCGCGAACTCGGGCAGCATCAATACGGGCGGTTTCGAC

TCGGGCGACCGGAACACCGGGTTTGGAAGCTCGGTCGACCAATCCGTTTCGAGCTCGGGCTTCGGC AACACCGGCATGAA

CCAACTCCGGCGGCTTCAACGTCGGCTTCTATAACTCGGGTGCCGGCACCGTGGGCATCGCAAACT CTGGCCTGCAGACC

ACAGGCATTGCGAACTCGGGCACCCTCAACACGGGTGTGCCGAACACGGGTGACCACAGCTCGGGGGGGCTTCAATCAGGG

CAGTGACCAGTCGGGCTTCTTCGGTCAGCCCTAA

FIG. 3.

SUBSTITUTE SHEET (RULE 26)

FIG. 6.

atyttgatcg	gegtäätääe	dadgdeoggt	gtgaagtgct	gttggccgtg	100
atgtoggstt	acagtctcgg	ogtgoocgac	gagacaggcc	ttggtgctga	150
cgcggcgcgc	gcgtgaagtg	gogotgacac	agcacattgg	ggtatccgcg	200
gagaccgatc	gggccgtcgt	coccaagong	cgccaggcct	atgacagoot	250
ggtgtgcggt	egeegeegge	ttggcgccat	tggagccgag	atcqaqaacq	300
oggtggccca	teagegegeg	argggeattg	acaccccggc	cagtacccat	350
sacttotoco	ggtttetege	caecaaagea	caccaccatca	cgcgagtgct	400
ggcagcaacc	gccgcggaat	cocaggeegg	cqcqqcqcqq	ttgcgatecc	450
tggcttcgtc	ctatcagget	gtgggatttg	gccccaaacc	ccaqqaqccq	500
octooggato	cagtgccatt	toogcoctac	cagoogaagg	tgtggggggc	550
gtaccagaca	cgtggccaag	acodggacsa	ggtogtoagg	acqttccatc	500
*appopogat	gagegegaga	ttccqctcqc	ttactogtgs	ccaattsgga	650
cotgatatog	coatggoott	gtogt			678

gatoogaatt oggcaogagt ogaggcdadd gottocatgg coaggccaog

FIG. 5.

ggatccgaat	totgcacgag	ggkygacgac	gamctttgca	cacgagegat	50
ggcaacoctc	acgloogcgo	aaaccccgcg	cgaggccgta	gagcastteg	100
togagotgat	ggtcgacgat	ocggtgogog	ggcgcgtgct	gttgctggcg	150
ccggcggtag	aaccggccct	gaccoggtog	ggcgcggagt	ggatgccsa	200
cttcatcgag	ttgctgcaac	gcaagttgto	ocquatcgtt	gatecagtte	250
tgcagaaact	ggtcgccacc	agcttgateg	gegetettae	aggtatgtta	300
accgcatato	tgaacggacg	gotgggagec	accogcaage	aattoatoga	350
ctactgcgtc	aucatgitge	toagcacege	cgcacctacg	caccgcaccg	400
ogagogggga	gáatccgaac	a			421
		-			

FIG. 4.

GAGTYGIANSGLQT TGIANSGTLNTGVANTGDHSSGGFNQGSDQSGFFGQP,

AGDTHTGFENAGFFAMGIGNAGNEDMGVGNGGSFNVGVGNAGNQSVGFGNAGTLNVGFANAGSINT GFANSGSINTGGFD SGURNTGPOSSVDQSVGSSGFGNTGMNSSGFFNTGNVBAGYGNNGDVQSGINNTNSGGFNVGFYNS

 $\label{lem:constraint} {\tt CNNLGLGNNGNNLIGFGLTGDNLVGLGALMSGLGNLGFGNSGNNNLIGFFNSGNNNVGFFNSGNNNFGGGNAGDINTGFGN\\$

Chilhranassand Athlesenighthlesenaghanlesenrefenlenenfusenlesentestnfegenlesfnles Gnieseniefenne

YEOMWAAOVAAMYG YHGGASAAAAQLISENSIGLOQALPAAFSALAAAIGLGNIGVGNLGGGNTGDYNLGSGNSGNANYGS GASGAANYSGNDG

mehrhemmysvippeinslpmftgagsapmlaasvavdglaaelavaassfgsvtsglaggswog aaaaamaaaapya gwlaaaaraagasagkavasafeaaraatvhpmlvaanrnafvolvlsnlfgonapaiaaaeam

>aTCC#3-His.pro

tgatcggtca	atgogoagta	ctggtgacot	agogoogoog	cggtggccat	50
catctcctcg	atcggcgcgg	accegteega	coagttogaa	tgcagatgca	100
gateccegeg	caatgoggca	aggategece	ctocacogag	atcotcagog	150
tcagogogta	attcagccag		bogoggdoag	accaggootg	200
ggcgatgact	ttogaggttt	tgggaccgat	accogcoage	gactgccage	250
tgttggcotg	geogtgooge	tgaage 276			
		FI	G. 7.		
ggatcogaat	totgoacgag	gangaagtca	tactgccgtc	atacacnttt	50
gtctytaccg		ogtgttgcgc	ggtggtgtgc	cagtctttqt	100
ogataggogg	cccgacacge	tcaacattga	tgasactcgc	atogtagaog	150
ccatcacccc	gcgaaccaag	gocatogtoc	cogttcacta	tgccggcgtg	200
acccacasass	tggacgcgat	catgaagatc	gccacgcace	acaacotgge	250
ggtggtcgaa	gaegeggeee	aaggogogat	ggcgtcgtat	agtgggggg	300
egeteggeag	catoggogac	chaggagege	totcatttoa	cgagaccaag	350
aatgtgattt	ocggcgaagg	cggcgccctg	cttgtcaact	cataagactt	400
cctgctccgg	gcagakatto	tcagggaaaa	gggcaccaat	mrcagoenge	450
teactt					456
		FI	G. 8.		
gatatoggat	cggaattegg	cáccaggtge	contaggaga	aceactggtq	50
cacaagaggt	togtoagtoc		gtatagggac	aggittecte	100
aagtttotga	ogogogogge		cgaactgtot	cacgacgttc	150
tasacccago	togogtgoog		cgaacagece	aaccettggg	200
acctgctcca	gccccaggat		gacatogagg	tgccasacca	250
tecegtegat	atggactett		agoctgttat	coccagagta	300
octtttatco	gttgagegae		ctegggggtg	c	341
			G. 9.		
gatccgaatt	cagagoggog	accogtgoto	caagctcctt	cagogtogto	50
acgggctcat	cctatecgge	agateageag	gaggtteate	ogcasagtgc	100
ggctgcaacc	tacogaotto	gtgegeggeg	aggaacgcgc	cocctagggg	150
tatecgeccg	cgtcagacaa	cagtgoctog	gtotgatogg	taataggcga	200
ccdectcdad	gtocacatco	gccacctgct	cgaaacgtca	ggtcttgggg	250
tgcggggtgt	accggacggt	atgogoccag	ategigeegt	ctcggaatac	300
gaaagtatog	actocytogt	ogactoggot	gaccgcggaa	ttcgcggtcc	350
actocaggaa	cagtatgtog	ccctcgeaga	tttgggtatt	tsagto	396
		FIG	i. 10.		
ggatcogaat	toggoacgag	gagtatoago	agaggtegga	gaaggtgotg	50
accgaataca	acaacaaggc				100
teaccaagea		accedecece	godtoogcaa	gagcagggat	150
					16 49 15

FIG. 11.

Egatcootgg cttootgatg co

ggatccgast	Loggoacgag	ccagaacoto	gookgecoog	ggoggcagng	50
acaccaactg	gscaccacgc	cgcggatcgg	cmgagcagcg	CC	92
		FIC	3. 12.		
gatoogaatt	oggcaogaga	agaatntgac	connuncting	tggotgatgo	50
gagagettne	tinttatte	ecoccantgg	ttggacgggg	togtcacase	100
gggcattota	agtoocyagg	gccacaeaag	gragtgrege	qqaacttett	150
ggcccaaacg	ggcaccoggc	tacgtgcgcs	cogagecegt	cgacaactgg	200
toggogagac	ggtccgggga	atccaccatc	gagaacgtcc	gtgctbcctc	250
gattacctcy	aascgggcgc	gogggatggt	cgcggcgagc	cgttgaccgt	300
totogagtgo	gaagaacacg	teatocgoog	aceacgcgat	gagogooggo	350
ttgtcgaatt	caygeageeg	ggoggogact	goggtggtga	ctteggtgag	400
cagogatage	gagagetgae	goaggtotto	ggcgatggcc	gggttggata	450
gagaaggsag	aacccaggcc	cgggtgagat	ggtcgatgtt	gtggtgegae	500
saaccggcat	acgcgcggtt	tacgcgcggc	cagtaccage	atcacctgga	550
togoggooog	gaacagggtg	googatttog	cggncaggat	cacotgnttt	600
gaggategg					609
		FIG	i. 13.		
ggatoogaat	toggcacgag	tgcggtgcct	atctgcgttg	gocagtacet	50
ogoggacotg	gogagtgcgg	acgcgcaygc	tatogaagtg	ggoctaaaga	100
sggoggacgt	ggogocogtt	gccgtscgsc	ctgcagcggc	geogeogtig	150
cgtgagtctg	cogoggtgag	accggaggoc	aggetgglgt	oggoggtggc	200
gccagctccc	gegggeaegt	cggcgtcggt	gctggottcg	gatcggggtg	250
coggogtatt :	ggggtttgcc	gggacogetg	gcaaggant.c	onttgggogt	300
		FIG	. 14.		
jąctąctącą	cgcactcgcg	ggtotgotgg	acqaqtqqac	googgtgate	50
lecadedced	aactgggcga	gcacccctac	acgccgatca	cgccggagto	100
gatooggogg	geogegeage	teggegaega	cctaceggtg	gogtggmage	150
reegeagega	gogotacaco	gagaagctgg	ccaccccga	caccagcgte	200
sccgacetgg	toggogacyt	cgacccgatc	aaggtigccg	agggccgcag	250
ctcggggat	C				261
		EIC	45		

PREDICTED PROTEIN SEQUENCE (SEC ID NO: 161)

VRHHEGHVAADDDCPQCASFCALTGY1EDIAENQRNAHHQKMRHGRCVEEVHLPVDVGEPRQPTGA
VADQDHRITFVFAHKHTPFRVCQDWHRQPFHGRADQHLGIDAFLCAACGVLLVDGVQRHFQRHG
PGERFGFPRVVVAGCIBQARVEVERFGGVPERAHGVGGNNRVATORLTDRHPIDRGLGEERFSV
GGQIDRERDQPQAIFAGXHVTFHCPQPRALHLVLTSRRHVERQRHRAEEDCHEVHAGPLGGASQSQG
HPGASEPPRHTHPSSFHGGGAAAGQQSDVHPFANLIAVDDRRAERRODEERGSAVQGKOPRCDEAD
PVADQQHPGDGADQCRPADPEHHDRHQRHQDHTQQGAGEPPRAESVTEDGLFDRDQLLTDRRVHQ
AVBGVVFHPMVVQLLFGLGCVMLLVEDGGASIGORAQVGEFGHBGQGRDGAGHDPAA

NECLEOTIDE SEQUENCE (SEQ 1D NG: 160)

TGAGATTGGCAGACCGGTGAGCACCGGGATACAGCCACGCAAAGTTCGTCACCACGAGGGCCACGTA GCAGCAGACGATCAGCCCCAGTGTGCGTCGTTCGGAGCCCTGACCGGGGTGATAKAKKATATC GCCGAGAACCAGCGAAATGCCCATCACCAGAAATGGCGCCATGUTUGCTGUGTAGAAGAAGAAGTACAT CTGCCGGTCGATGTCGGCGAACCACGGCAGCCAACCGGCGCAGTAGCCGACCAGGACCACCGCATA CATCGCGGGCGTGCCGACCACCATCTCGGCCTTGACGCACGACTGTGCGCCGCCGCAGCCTGCAACCTC TTGCTGGTCGATGGCGTACAGCACCGGCCGCAACGACATGGGCCAGGTCCACGGTTTGGATTCCCA AGGGTGGTAGTTGCCTGCGGAATTCGTCAGGCCCGCGTGGAAGTGGAACGCTTTGGCGGTGTAGTG ATGCCGATCGCTGTCTCGGACGCGAACCACGGGGCGTAGGTGGCCAGATAGACCGCGAACGG GATCAACCCCAGCGCATACCCGCTGGGAAGCACGTCACGCCGCCACTGTCCCCAGCCACGGTCTTTG CACTTGGTACTGACGTCGCGCCGCCGCCGCGCGCGCGCCGCCGCACACACCACCCAA GUGUACACUTACCUGUGGTCCCCACACGGTGGUGGCGCTGCGGCCAGCAGCAGCAGCATGTGCAT COSTTOGOGAROCTGATOGOGGTOGACGATGAGOGCGCGAACGCOGGACGAGGAACGTOAG GAAGCIGTCCAGCAGCGCGCCGCGCGCGCGAAGCTGACCCCGTCCCAGATCAGCAGCACCCC GGCGATGGCGCCGACCAATGTCGACCGGCTGATCCGCCGCACGATCCGCCACCAGCGCCCACCAG GACCACACCAGCAGGGCGCGGTGAACCGCCAGCCGAATCCGTTGTAACCGAAGATGGCCTCCCC GATCGCGATCAGCTGCTTACCGACCGGCGGTGAACCACCAGGCCGTACCGGGGTTGTCTTCCAC CCCATGGTTGTTCAGCACCTGCCAGGCCTGGGGTGCGTAATGCTTCTCGTCGAAGATGGGGGTTGCC GGCATCGGTCAGCGAGCCCAGGTTCAGGAACCGGGTCACCGTGGCCAGCGTGATCAGGCCGGT CACGATCCAGCCGCGTAA

NOTES: UNKNOWN PROTEIN FROM COSMID MTC1237

FIG. 16.

10711

MO-2 PREDICTED PROTEIN SEQUENCE (SEO ID NO: 163)

VALVVOKYGGSSVADAERIRRJÆRIVATKKOGNDVVVVVSAMODTIDDLLDLAQQVCPAPPPREL EMILITAGERISNALVAMAIESLGAHARSFICSOGGVITTGENAKTIDVTPARLOTALEEGRVVL VAGFQGVSQDIKDVITLGRGGSDITAVAMAALGADVCEIYIDVEDIFSADPRIVNNASKLDTVIF EEMLEMAACGAKVLMIKCVEYARRHNIDVHVRSSYSDRPGTVVVGSIKDVPMEDPILITGVAHDRSE AKVITUGLPDIFGYVAKVFRAVADADVNIDMVLOMVSKVEKGKTDIFFTCSRDVGPAAVEKLDSLR NEIGFSQLLYDDHIGKVSLIGAGMRSHPGVTATFCEALAAVGVNIELISTSEIRISVLCRDTELDK AVVALHERFGLEGDEERTVYASTGR

NUCLEOTIDE SEQUENCE (SEQ ID NO. 162)

GCCGAACGCATCGTCGCCACCAAGAAGCAAGGCAATGACGTCGTCGTCGTCTCCTCTGCCATGGGGGA TACCACCGACGACCTGCTGGATCTGGCTCAGCAGGTGTGCCCGGCGCGCCGCCTCGGGAGCTGGA CATGCTGCTTACCGCCGGTGAACGCATCTCGAATGCGTTGGTGGCCATGGCCATCGAGTCGCTCGG COCOCATGCCCGGTCGTTCACCGGTTCGCAGGCCGGGGTGATCACCACGCGCACCCACGGCAACGC GOOGGATTOCAAGGGGTCAGCCAGGACACCAAGGATGTCACGACGTTGGGCCGCGGCGGCTCGGA CACCACCCCCTCGCCATGGCCGCCCCCCTGGGTGCCGATGTCTGTGAGATCTACACCGACGTGGA COGCATOTTCAGCGCCGACCCGCGCATCGTGCGCAACGCCCGAAAGCTCGACACCGTGACCTTCGA GGAAATGCTCGAGATGCCGGCCTGCGGCGCCAAGGTGCTGATGCTGCGTTGCGTGGAATACGCTCG CCGCCATAATATTCCGGTGCACGTCGGTCGTCGTACTCGGACAGACCGGGCACCGTCGTTGTCGC ATTGATCAAGGACGTACCCATGGAAGACCCCATCCTGACCGGAGTCGCGCACGACCGCAGGAGGC CAAGGTGACCATCGTCGGCTGCCCGACATCCCGGGTATGCGGCCCAAGGTGTTTAGGGCGGTGGG CGACGCCGACGTCAACATCGACATGGTGCTGCAGAACGTCTCCAAGGTCGAGGCCGAGACGGCAAGACCGA CHTCACCTTCACCTGCTCCCGCGACGTCGGGCCCGCCGCCGGGAAAAACTGGACTCGCTCAGAAA CGAGATOGGCTTCTCACAGCTGCTGTACGACGACCACATOGGCAAGGTATCGCTGATCGCTGCCGG CATGCCCACCCCCGGGTCACCCCGACCTTCTGTGAGGCGCTGGCGGCGGTGGGGGTCAACAT CGAGCTGATCTCCACCTCGGAGATCAGGATCTCGGTGTTGTGCCGCGACACCGAACTGGACAAGGC CETGGTCGCGCTGCATGAAGCGTTCGGGCTCGGCGGCGAGGAGGCCACGGTGTACGCGGGGAC gggacggtagatgggctgtcaatagggatcgtgggggccaccggtcaggtgaggtcatgcg CACGTTGCTCGACGAGCGGGATTTCCCGGCGAGGGCGGTGCGGTTCTTCGCGTCGGCCCGATCGCA GGGCCGCAAGCTGGCCTTCCGCGGCCAGGAGATCGAAGTGGAAGACGCCGAGACGGCCGACLCGAG CGGCTGGATATCGCGTTCTTCTCCGCCGGCTCGGCCATGTCGAAGGTGCAGGCGCCCCCTTTGC ggracergagtcacgutgatcgacaactcgtcggcgtggcgtaaggaccccgacgtgccgttggt ggtgtccgaggtgaactttgaacgggacgggcaccgccggccCaaggctcctgccgctcgtgccga CCSTCCACCAGGTGGTAGGAGCGAACGAAGATTCCACCGTCGTCGTCAACGTGGCCGCATTGCCG TACGAATCGACGACGCTGAGGTGGCTGGTGCCATGCTCAGGCACTGGCGGGGGGGACGGCCGTCGGT GCGCCGAAGTCCC

NOTES: M.tb aspartokinase

FIG. 17.

>Full-length TbH4/XP-1 (MTB48) Open Reading Frame (SEO ID NO: 164)

atgacgcactcgcagaccgtgacggtggatcagcaagagattttgaacagggccaacgaggtggag GCCCGATGCCGACCCACCGACTGATGTCCCCATCACACCGTGCGAACTCACGGCGGCTAAAAAC GCCGCCCAACAGCTGGTATTGTCCGCCGACAGATGCGGGAATACCTGGCGGCCGGTGCCAAAGAG CCGCASCCTCTGCCGACCTCGCTGCGCCAACGCGCCAAGGCGTATGGCGAGGTTGATGAGGAGGCT ACACTTCGCCCGAACTAACCGATACGCCGAGGGTGGCCACGGCCGGTGAACCCAACTTCATGGATC PCANAGÁGGGGGAAGGAAGCTCGARACGGGGGACCAAGGCGCATCGCTUGCGCACTTTGCGGATG GUTGGAACACTTTCAACCTGACCTGCRAGGCGACGTCAAGCGGTTCCGGGGGTTTGACAACTGGG AAGGCGATGCGCTACCGCTTGCGAGGCTTCGCTCGATCAACACGGCAATGGATACTCCACATGG CCAAATTGAGCGCTGCGATGGCCAAGCAGGCTCAATATGTCGCGCAGCTGCACGTGTGGGCTAGGC gggaacatccgacttatsaagacatastcgggctcgaacggctttacgcggaaaacccttcggccc CCCACCAAATTCTCCCGGTGTACGCGGAGTATCAGCAGAGGTCGGAGAAGGTGCTGACCGAATACA ACAACAAGGCAGCCCTGGAACCGGTAAACCGGCGAAGCCTCCCCCGCCATCAAGATCGACCCGC COCCCCCCCCAGAGCAGGGATTGATCCCTGGCTTCCTGATGCCGCCGTCTGACGCCTCCGGTG TGACTCCCGGTACCGGGATGCCAGCCGCACCGATGCTTCCGCCTACCGGATCGCCGGGTGGTGGCC TGGCGCTCAAAGCGGCATCGCTCGGTGGCGGTGGAGGCGGGGGGGTGCCGTCGGCGCCGTTGGGAT CCGCGATCGGGGCGCCGAATCGGTGCGGCCCGCTGGCGCTGGTGACATTGCCGGCTTAGGCCAGG annana agregoroccaagteraaggettétcagcaggaagacgaggegetétabaccgaggate GGCATGGACCGAGGCGTCATTGGTAACCGTCGGCGCCAGGACAGTAAGGAGTCGAAG

FIG. 18.

SEQUENCE LISTING

	(1) GENE	RAL INFORMATION:
5	256	to purply 12 years areas
	(3.)	APPLICANT: (A) NAME: Corixa Corporation
		(B) STREET: Suite 200, 1124 Columbia Street
		(C) CITY: Sentile
10		(D) STATE: Washington
10		(E) COUNTRY: USA
		(P) POSTAL CODE (NIP) + 98104
		(G) TRLEPHONE: (206) 754-5830
		(N) TELEFAX: (206) 754-5394
15		(I) TELEX:
		7.00.3 **********************************
	(\$1)	TITLE OF INVENTION: Compounds for Immunotherapy and Diagnosis of Tuberculosis and Methods of Their Use
20	74443	NUMBER OF SEQUENCES: 144
2010	10000	THE THE PARTY OF THE PARTY WAT
	(34)	COPRESPONDENCE ANDRESS:
		(A) ADDRESSE: Townsend and Townsend and Crew LLP
		(B) STREET: Two Embarcadero Center, Eighth Floor
25		(C) CITY: San Francisco
		(D) STATE: California
		(B) COUSTRY: USA
		(F) EIP: 94111-3839
30	(3)	COMPUTER READABLE FORM:
		(A) MEDIUM TYPS: Diskette
		(B) COMPUTER: IBM compatible
		(C) OPERATING SYSTEM: Windows
35		(D) SOFTWARE: FastSEQ for Windows Version 2.0b
20	Inch N	CURRENT APPLICATION DATA:
	10.27	(A) APPLICATION NUMBER: PCT/US98/15407
		(B) FILING DATE: 20-HAY-1998
		(C) CLASSIFICATION:
40		
	(vii)	PRIOR APPLICATION DATA:
		(A) APPLICATION NUMBER: US 00/859,381
		(B) FILING DATE: 20-MAY-1997
		the Contract
45		(A) APPLICATION MUMBER: US 05/073,010
		(B) FILING DATE: 05-MAY-1998
	toxid it is	ATTORNEY/AGENT INFORMATION:
	247773	(A) NAME: Bastian, Kevin L.
50		(B) REGISTRATION NUMBER: 34.774
-		(C) REFERENCE/DOCKET NUMBER: 14058-87-1PC
		The second secon
	(ix)	TELECOMMUNICATION INFORMATION:
		(A) TBLEPHONE: (415) 576-0200
55		(B) TELEFAX: (415) 576-0300
		(C) TELEX:

```
(2) INFORMATION FOR SEC ID NO:1:
          (i) SEQUENCE CHARACTERISTICS:
               (A) LENGTH: 1886 base pairs
 5
               (B) TYPE: nucleic acid
               (C) STRANDEDMESS: single
               (D) TOPOLOGY: linear
         (11) MOLECULE TYPE: DNA (genomic)
16
         (xi) SECURNCE DESCRIPTION: SEC ID NO:1:
     CGCTCTGGTG ACCACCAACT TCTTCGGTGT CAACACCATU CCGATCGCCC TCAACGAGGC
     COACTACTS COCATOTOGA TOURSECOSC CACCOTCATO AGCCACTATC AAGCCOTCOC
                                                                          120
15
     GCACGAAATC TOGTGTCYCC AYGAATANGC CAGTYCGGGA AAGCCGTYGG CCAGTATCAC
     CACGOSTOCS COOGOCTCAC COSCCTCGAC CACTCGCAGT CGCACGCCGT TGGTATCAAC
                                                                         240
     TAACCGINCE STANGIGGC CCATCGICTC ACCARATORC ACCGGGCACC GGCCTGAGAA
     GOSCITISGGS AGCANCCAGA SGCGATIGIC GCGGGTSCTG CCGCGCATCA TIGATCOSCC
     GECCHACCA NTONGGECTE CETTEACNTE CHEATUNEAC TYPETNICA GETGGCATON
20 CTACAGCTCA CAGTGACTOC CCCACGATTO COMICCAGGT CCAGTTCAAA TTCCGGTGAA
   TYCGCCGACA AAAGCAGCAG GYCAACCAAC CGCAGTCAGT CGAGGGTOUU AAACGTGAGC
     CARTOSOTIS AATSSCTTK: TSCADISACA COMSTCACES SCTTASCOGA CASCAGOSIA 600
     ATAGCTCAGG CUCGCCTATAG AGTCCTATAG AAACATITUC TGATAGAATT AACCGCCCTCTC
    TEGGCOTGAT CTTGATACGG CTCGCCGTGC GACCGGTTGG CTCAGTAGCT GACCACCATG
     TARLECATEC TEGGERGOTS TETRETARGS CGRGACACUS CATTEFFGGG SCTSCATCSC
     AAATCHFTCX GAGGATGTAG CACTGCCGTT ATCCCGGAT ACCAAACCAC CCGGAACCAG
     OGCTATUCCA STORCTOTOL GAUGGAGGCU OTTTOGCTTT COGTTGCCGG ATRACTCCCG
     AGFRIGATATE GUCUPPATCA NATICAGGET ITTETTEGCA AGGRACEGGI GITCGCTATA 960
     TTCGGATATC TCGGACGGAT AATTACTAAA ACTTCAGTGG TTTAGATAAG GCCXCCXCAA 1020
30 TACTINICCO ATCTTOCCOA GCOCAACOGA TITCCATCOT COOTITICOT COCCITATCA 1080
     AMCAPGATON GAGATAATGA CAGATCEGOO TAGCTAGOTG TTTAGCGGAC GCGATTTAGG 1140
     ACRACOGAGA TITOCTTTGC CTCGCRACCA TGAGAGUGCC CCGCTTCGAC GCCGAATCXX 1200
     STREETSATE STORSTTAGE ACAGCUCTGA TYPEGCUACU SECSAGGIVA TISTECCUSC
     CACCAGGEOG COGCOGCTA GOOCCATGAG CACCATAT AGACTOTOCT GCAACAGATC 1320
35 TCATACCCAT CGAAGGCGAA GCGCAGGCAT CGACGTCGGA GACACTGCCT TGGGATCGGG 1185
     CCGCCTACAC GGCGGTTGGC GCATTGTCGC ACCGCAGTTG CAGGAGGGCA AATGTGCGCA 1440
     GACGATGTAG TOGACAACAA GTGNACATOC COFCTTCACG AACTCAAAAC TGACGATGTG 1500
     CTTABCATGE AAAAAACTGT TGACATGGGC CAAGCATGAC AGCCAGACTG TAGACCTACG 1960
     CCTGCAATDC AGAACCAAGG HTATGCATGG AATCGAGGAC CUTTGAGATA GGCGGCAGGC
    ATGRACCAGAG COTTCATCAT CGATCGAACG ATCAGTRCCA TEGACGGCTT GTACGACCTT 1680
46
     CIGGGGATTG GAATACOCKA CCAAGGGGGT ATCOTTEACT COTCACTAGA GTACTICGAA 1740
AAAGGCCOTGG AGGAGCTGGC AGCAGCGTTT CCGGGTGATG GCTGGTTAGG TTCUCCCGCC 1860
     GACAAATACO CCCCCAAAAA CCGCAACCAC GTCAATTTTT TCCAGGAACT GCCAGACCTC 1866
     GATOGTCAGC TCATCAGCCT GATCCA
                                                                         1886
45
     (2) IMPORMATION FOR SEQ ID NO:2:
          (i) REQUENCE CHARACTERISTICS (
               (A) LENGTH: 2305 bass pairs
50
               (B) TYPE: nucleic acid
               (C) STRANDEDNESS: single
               (D) TOPOLOGY: Linear
         (ii) MOLECULE TYPE: DNA (senomic)
55
         (xi) SEQUENCE DESCRIPTION: SEC ID NO:2:
    GCCACGCGCT GGCCGCGCAR TACACCGARA TTGCCARCGGA ACTOGCRAGC GTGCTCGCTG
```

	CONTROCAGGC	AAGCTCGTGG	CAGGGGCCCA	GCGCCGACCG	GTTCGTCGTC	GCCCATCAAC	120
	OGTTCCGGTA	TIGGCTAACC	CACGCTGCCA	CGCTGGCCAC	CGCAGCAGCC	GCCGCGCACN	180
	AAACGGCCCGC	COCCOGGTAT	ACCTCCGCAT	TOCOGGGGGAT	OCCTACGCTA	GCCGAGTTGG	240
	COCCAACCA	TUCCATGCAC	GGCGCTCTGG	TOACCACCAA	CTTCTTCGGT	GTCAACACCA	300
5	TOTOGGATOGG	CCTCAACGAG	GCCGACTACC	TOCOCATORS	GATCCAGGCC	OCCACCOTCA	360
	TORGCCACTA	TCARGCCGTC	GCGCACGAAA	GCGTEGCGGC	GACCCCCAGC	acceccecco	420
	COCCOCAGAT	AGTGACCAGT	GOGGCCAGCT	CGGCGGCTAG	CAGCAGCTTC	CCCGACCCGA	480
	CCAAATTGAT	COTGCAGCTA	CTCAAGGATT	TOCTOGAGCT	GCTGCGCTAT	CTGGCTGTTG	540
	AGCTGCTGCC	QQQQCQQCTC	GGCGACCTCA	TCOCCCAGGT	GTTGGACTGG	TTCATCTCUT	600
10	10010100000	TOCAGTOTTC	ACGITTCICG	CCTACCTGGT	GCTGGACCCA	CTGATCTATT	660
	TOGGACCGTT	COCCCCCCTG	ACGAGTCCGG	TCCTGTTGCC	TGCTGTGGAG	TTACGCAACC	720
	GCCTCAAAAC	CGCCACCGGA	CTGACGCTGC	CACCTACCGT	CATTITCGAT	CATCCCACTC	780
	CCACTGCGGT	COCCGAGTAT	GTCGCCCAGC	AAATSTCTGG	CAGCCGCCCA	ACGGAATCCG	940
	GEGATOCGAC	GTCGCAGGTT	OTCGAACCCG	CTCGTGCCGA	ATTOGGCACG	AGTGCTGTTC	900
15	ATCAAATCCC	CCCGAGACCT	GCGGACACCC	GGCGCGCTTG	COGACATOGA	GATGATGTCC	968
	COCGAGATAG	CAGAATTOCC	CAACATOGTG	ATCCTCCCCC	OCTIVACCOU	ACCGAACGGG	3030
	GAACCTCTGA	AGGAGRCCAA	GGTCTCGTTT	CAGGCTGGTG	AAGTGGGCGG	CAAGCTUGAC	1080
	GAAGCGACCA	CCCTGCTOGA	AGAGCACGGA	GGCGAGCTGG	ACCAGCTGAC	0280393369	1140
	CACCAGTTUG	CCGACGCCCT	CGCCCAAATA	COCAACGAAA	TCAATGGGGC	COTGOCCAGC	1200
20	TOGAGOGGGA	TAGTCAACAC	CCTGCAGGCC	ATCATEGACC	TGATGOGCGG	TGACAAGACC	1260
	ATCCGACAAC	TUGARARATEC	GTCCCAATAT	GTCGGGCGCA	TGCGGGCTCT	GGGGGACAAT	1320
	CTGAGCGGGA	CCGTCACCGA	TGCCGAACAA	ATCGCCACTT	GGGCCAGCCC	TATGGTCAAC	1380
	GCCCTCAACT	CCAGCCCGGT	OTOTAACAGC	GATCCCCCCT	GTCGGACGTC	GCGCGCACAG	3440
	TTGGCGGGGA	TTOTCCAGGC	GCAGGACGAC	GOCCTGCTCA	GOTCCATCAG	AGCGCTAGCC	1.500
25	GECACCUTUC	AACAGACGCA	GGAATACCAG	ACACTOGOCO	GRACGGTGAG	CACACTGGAC	1560
	GOGCAACTGA	AGCAAGTCGT	CAGCACCCTC	AAAGCGGTCG	ACGGCCTACC	CACCARATTO	1628
	CCTCAAATGC	AGCAAGGAGC	CARCGCTCTC	GCCGACGGCA	GCGCAGCGCT	GOCGGCAGGC	1689
	GYGCAGGART	TOGTCGATCA	GGTCAAAAAG	ATGGGCTCAG	COCTORACGA	GGCCGCCGAC	1740
	TICCICTECKS	GGRATCARGO	CGATCCGGAC	AAGCOSTCAA	TOGCOGOCTT	CARCATTOCA	1800
30	CCGCAGATTT	TTTCGAGGGA	CGAGTTCAAG	AAGGGCGCCC	AGATTTTCCT	GTCGGCCGAT	1860
	GGTCATGCGG	CGCGGTACTT	COTGCAGAGC	GCGCTGAATC	COGCCACCAC	CEAGGCGATG	1.920
					GACCGAATAC		1980
					COGATATORS		2040
	AACAGCGATA	TGAAATTCAT	COTCATTON	ACCATCGTTA	TOTTATTCTT	GATTCTCOTC	2300
35	ATPCTONTEC	GCGCACTTGT	GGNTCCGATA	TATCTGATAG	GCTCGGTGCT	GATTTCTTAC	2160
					TACTGGGCCA		5330
				TIGGTIGCCA	TORGOGCTGA	CTACAACATG	2280
	CTOCTCATTT	CAOSCATOOS	CGACG				2305
40	(2) IMPORM	TION FOR SI	NO ID NO:31				
		C. W. S.					
			RACTERISTIC				
			3742 base j				
45			scleic acid				
40		(C) MIKANDE	ONESS: sing				
		(ii) TOPCILOGS	es tanger				
	(ii) 300	LECULE TYPI	: DNA (geno	omic)			
30	(xi) 88	QUENCE DES	CRIPTION: SI	BQ ID NO:3:			
	cicio con com como	TO S NO CONTROL OF	x x cmm, cacamo	anameran	COGCGCGTGC	a management of	60
					AGAGCGGATC		120
					TGCGGAAGAT		180
55					GCGCCCCGGGC		246
44.44	xx-2000000	MAS YOUNG THE	SAMMAN SAMMAN	03 0 - 10 Cale 19 19 19 19 19 19 19 19 19 19 19 19 19	MANAGER CONTRACT	CASAL PASSAGE TANA	46.5 6.

GCALOSCICO GCTCCOSCAC CCGGCCCCG GCTGCCAACA CCCCACOATT GAGAIGGAAG CCGATCACCC GTGCCATGAC ATCAGCCGAC GCTCGATAGT ACGGCCCCC GACACUGGC

AGATCATOCT TOAGCTCGOC CAGCCGOCGG TOMOTOCCGA ACAGCGCCAG CGGCGTRAAC

366

420

	CONDAGGCUA.	GCATGCGCTG	CACCACCAGC	ACACCCTCGG	CGATCACCAA	CUCCTTOCCG	480
	GTCGGCAGAT	COGGACNACN	GTOGATGCTG	TECAGOTCAC	GGAAATCGTC	GAGCCOTGGG	540
	TOTTOGOGAT	CGCAGACGTC	CTGAACATCG	AGGUCGTCGG	derecresed	ACAACGGCCT	600
	TCGGTCAC9G	GCTTTCGTCG	ACCAGAGCCA	GCATCAGATC	SGCGGCGCTG	OGCAGGATGT	660
5	CACGCTCGCT	GOGGTTCAGC	GTCGCGAGCC	GCTCAGCCAG	CCACTCTTGC	AGAGAGCCOT	720
	TGCTGGGATT	AATTGGGAGA	GGAAGACAGC	ATGTCGTTCG	TGACCACACA	GCCGGAAGCC	780
	CTOGCAGCTG	COGCGGGGAA	CCTACAGGGT	AFTGGCACGA	CAATGAACGC	CCAGAACGCG	840
		CTCCARCCAC					900
	ACCGCGGCTC	ASTITUCTEC	OCAC909CAG	ATGTACCAAA	COGTCAGCCC	CCAGGCCGCG	260
10	CCCATTCACG	AAATOTTCOT	GAACACGCYG	OTOOCCACTT.	CTOGCTCATA	CACCACCYCC	1020
	GAGGGGGCCA	ACGCAGCCCC	TGCCGGCTGA	ACQGGCTCGC	ACGAACCTGC	TGAAGGAGAH	1089
	GGGGAACATC	CGGAGTTCTC	GGGTCAGGGG	TTGCGCCAGC	GCCCAGCCGA	TTCAGNTATC	1140
	GGCGTCCATA	ACAGCAGACG	ATCTAGGCAT	TCAGTACTAA	GGAGACAGGC	AACATGGCCT	1200
	CACCETTTAT	GACGGATCCG	CATGOGATOC	GCCACATGGC	GUGCCOTTTT	GAGGTGCACG	1260
15	CCCAGACGGT	GGAGGACGAG	GCTCGCCGGA	TOTOGGCGTC	CCCCCAAAAC	ATTICCGOIG	1320
	CGGGCTGGAG	TOGCATOGCC	GAGGOGACCT	COCTAGAÇÃO	CATGACCTAG	ATGAATCAGG	1380
	CCTTTTCSCAA	CATCGTGAAC	ATGCTGCACO	SCOTOCOTCA	COGGCTGGTT	COCGACGCCA	1440
	ACRANTACCA	ACAGCANGAG	CAGGCCTCCC	AGCAGATCCT	GAGCAGNTAG	CGCCGAAAGC	1509
	CACAGCTUNG	TACGNTTTCF	CACATTAGGA	GAACACCAAT	ATTROCHATTA	ATTACCAGTT	1,580
20	COGGGGACGTC	GACGCTCATG	GCGCCATGAT	COSCSCTUAG	GCGGCGTCGC	TTOAGGCGGA	1520
						GOGOCGGTTC	1680
	GOTGGCTTGC	CAGGAGTTCA	TTACCCRGTT	GGGCCGTAAC	TTCCAGGTGA	TCTACGAGCA	1740
	QQ						1742
20.00							
25	(3) IMPORM	ATION FOR SI	8Q ID NO:4:				
	(i) si	EQUENCE CHAI	RACTERISTIC	S t			
		(A) LEMGTH:		pairs			
			auleic soid				
30			MESS: sing	Le			
		(D) TOPOLOG	7: linear				
	(3.3.) MS	OLECOLE TYPE	8: DNA (gen	omic)			
35	(xi) Si	EQUENCE DESC	ERIPTION: N	SQ ID NO:4:			

35	(x1) S	EQUENCE DES	CRIPTION: S	EQ ID NO:4:			
	GTTGATTOCG	Tregeogege	CGCCGAAGAC	CACCAACTCC	acressares	TOGGACAGGC	60
	ogrigcated	GTCAGCTGGC	CGAATCCCAA	TGATTOUTGO	CTCMGTGCGG	TTGCTGGGCT	3.20
	COATTACCC	CACOGARAGG	ACGACGATCO	TTOSTTTGCT	COGTCAGTEG	TACTTGGCGA	180
40	CHRISCATORC	GOOGTTTCTT	ACCTOGATOG	CACAGCAGCT	GACCTTCGGC	CCAGGGGGCA	249
	CAACGGCTGG	CTCCGGCGGA	GCCTGGTACC	CAACGCCACA	ATTOGCOGGC	CTGGGTGCAG	300
	gcccggcggr	GTCCGCGAGT	TTGGCGCGGG	CGGAGCCGGT	COGGAGGTTG	regardeese	356
	CAAGTTGGGC	COTOGGGGGG	COGGCCTTOG	COGAGAAGCC	TRAGGCGGGC	ACGCCGATGT	€20
	CCGTCATCGG	CURRECUTOC	AGCTGCGGTC	AGGGAGGCCT	GCTTCGAGGC	ATROCECTES	488
45	CGAGAGCGGG	GCGGCGTACA	eececcrrcs	CTCACCGATA	COGGTTCCCC	CACAGCGTGA	540
	TTACCCGGTC	TOCOTOGGGG	COATACCTTT	CONTOCUCTO	racocaacca	CCGGGAAATGC	600
	TGCAGATAGC	GATOGACCGC	SCC00TCGOT	BAACGCCGCA	CACGGCACTA	TCAATGCGCA	680
	COCKOGGCGT	TORTGCCAAA	TYGACCGTCC	CGACGGGGGCT	TTATCTGOGG	CAAGATTTCA	720
	TECCERGCCC	GOTCGGTGGG	COGATABATA	COCTOCTCAG	COCCACTOTY	CCGGCTGAAT	790
50	TCGATGCTCT	gggcgcccgc	TCGACGCCGA	GTATCTCGAG	TGGGCCGCAA	ACCCOGTCAA	840
	ACCCTOTTAC	TOTOCCOTTA	CCACAGGTGA	ATTTGCOGTO	CCAACTYGGTG	AACACTTROS	900
	AACGOGTGGC	ATCGALATCA	ACTTOTTGOO	THOUGAGEGAT	CTACTCTCTT	SCAGRGAGCC	960
	OTTGCTGGGA	TTAATTOGGA	GAGGAAGACA	GCATGTCGTT	CSTGACCACA	CAGCCGGAAG	1020
	CCCTOGCAGC	TGGGGCGGGG	BACCYACAGG	GTATTGGCAC	GACAATGAAC	GOCCAGAACG	1880
55	CGGCCGCGCGC	TOCTCCAACC	ACCOGAGTAG	TOCCCOCADE	COCCGATGAA	GTATCAGCTC	1140
	TGACCGCGGC	TCAGTTTGCT	GCGCACGCGC	AGATGTACCA	AACGGTCAGC	GCCCRGGCCG	1200
	CSGCCATTCA	CGASATGTTC	GTGAACACOC	TOGTGGCCAG	TICTGGCTCA	TACGOGGCCA	1260
	CCGAGGCGGC	CAACGCAGCC	GCTGCCQGCT	BAACGGGCTC	OCACURACCT	GCTGAAGGAG	1320

	AGGGGGAACA TOOGGAGTTO TOGGGTCA	GG GGTTGCGCCA	GCGCCCAGCC	GATTCAGCTA	1386
	TORGOGICCA TAACAGCAGA CGATCTAG				1440
	CTCACGTTTT ATGACGGATC CGCATGOS				1500
	COCCOMONICO CERGAGIGACIO AGGICECCIO				3560
5	TOCGOGCTOG ACTGGCATGG CCGAGGCS				1620
	GEOGRAFICGO AACATOONSA ACATOONS	CA COGCOTTACCT	GACGGGCTGG	TTCOURAGE	1680
	CAACAACTAC GAACAGCAAG AGCAGGCC	TO CCAGCAGATO	CTHACCAGCT	AGCGCCGAAA	1740
	GCCACAGCTG CGTACGCTTT CTCACATT				1,000
	TTUGGGGRCG TOWACGCTCA TOWCGCCA	TO ATCCGCGCTC	REGUGGGGGGG	OCTIGACION	1860
10	GAGCATCAGG CCATCGTTCG TGATGTGT				1920
	TOSSTERCTY SCCASSAGTY CATTACCC				1980
	CAGGCCAACG CCCACGGGCA GAAGGTGC				2049
	AGGGCCGTCG GCTCCAGCTG GGCCTAAA				2100
	OCCUPATION TONTOTONIC TOCACTTA				2160
15	CAACAGAGTA COOGCACOGA CATICACOG	TC AACRTCGACG	GCTTCTGGAT	CCTTCAGGCG	2220
	CTACTEGATA TOCGCCACCT TGCGCCTG	AG TTACGTTGCC	GOCCOTACGT	CTCCACCGAT	2280
	TOCAATGACT GGCTAAAGGA GCACCCCC				2340
	GTCAACGACG CGGTCRACGA ACAGGTCG				2400
	CTTGAAGTEG TEGECETTGET GTCACGCG				2450
30	AACCAGCCC CHGGTTCGCG TGACATCC				2520
M.V	CGAGGCCASC ACTGGCTGTC GGCGGTAC				
	ACCOTONO ATACCOCOTO CATORCO				
	CACGCGACC CAGCCGGGAT CAACGCGG				2700
	ATTCOGCAGE AGGCAGGAGG CGGTGTCG				2750
25	GCCGGUATCC TTGGCCGATCT CGTTGAGC				2820
Service .	CCATOGETTC TTCCCG	and conductional	cocooomicc	resoroners:	2836
30	(i) SEQUENCE CHARACTERIST (A) LENGTH: 906 base (B) TYPE: Ducleic ac (C) STRANDEDREES: si	pairs id			
	(D) TOPOLOGY: Linear				
35	70.7 CO.COM. CO.C. M. CO.C.				
	(ii) MIECULE TYPE: DEA (9	enomic)			
	(xi) SEQUENCE DESCRIPTION:	SEQ ID NO:5:			
40	AACATGCTGC ACGGGCTGCG TGACGGGC				60
	GAGCAGGCT CCCAGCAGAT CCTCAGCA				120
	ACAAGCGAAG GAGAACAGGT TOURTGAC	CA TCAACTATCA	GTTCGGTTGAT	GTCGACGCTC	1.80
	ACGGOGCCAT GATCCGCGCY CAGGCCGG	OT TUCTOGAGGC	COAACATCAS	CCCATCATTC	240
	GINATOTETT GACCGCGAGT GACTTYNG	GO 9009000000	TTCOOCGOCC	TOCCAGGOOT	390
45	TCATTACCCA ATTGGGCCGT AACTTCCA	GG TGATCTACGA	ACAGGCCAAC	GCCCACGGGC	360
	AGAAGGTGCA GGCTGCGGGC AACAACAT	GG CGCAAACCGA	CAGCGCCGTC	GGCTCCRGCT	420
	OGGCCTGACA CCAGGCCAAG GCCAGGGA	CC TGCTGTACGA	GTGAAGGTTC	CTCGCGTGAT	480
	CCTTCGGGTG GCRGTCTAGG TGGTCAGT	OC TODOGOTOTTG	GROOTETSCT	GCTTGGCGGG	540
	TRUTTOGGTG CTGGTCAGTG CTGCTCGG	GC TOGGGTGAGG	ACCTCGAGGC	CCAGGTAGGG	600
50	COSTCETTES ATCCATTEST CSTSTEST	TC GGCGAGGAOO	GCTCCGACGA	GGCGGATGAT	660
	CGAGGCGCGO TCGGGGAAGA TGCCCACG	AC STESSTICSS	COTCOTACCT	CTCGGTTGAG	720
	SCRTTCCTGG GGCTTGTTGG ACCAGATT				780
	CGCCAGCAGG TCGGTGCGGG CGGTGTCC				
	CAGAGCOTCG AGTACCCGAT CATATIGG				900
55					

(2) INFORMATION FOR SEQ ID NO:6:

(i) SEQUENCE CHARACTERISTICS:

(D) TOPOLOGY: linear (ii) MOLECULE TYPE: DEA (genomic) (xi) SEQUENCE DESCRIPTION: SEG ID NO:6: (xi) SEQUENCE SEG ID NO:6: (xi) SEQUENCE DESCRIPTION: SEG ID NO:6: (xi) SEQUENCE SEG ID NO:6: (xi) SEQUENCE DESCRIPTION: SEG ID NO:6: (xi) S		(A) LENTH: 1965 base pairs (8) TYPE: nucleic acid	
(ii) MOLECULE TYPE: DEA (genomic) (xi) SEQUENCE DESCRIPTION: SEQ ID NO:6: 30 OCTOCCOGA TOTOGGCCTC COCCEANAC ATTOCCOGT CONGENCOGA TOGGATOGCC ANGUEROCA COCCEANAC ATTOCCOGT COTTOGGCC CATOCCCCA ANGUEROCA CATOCCCCA CATOCCCCA CATOCCCCA CATOCCCCA CATOCCCCCA CATOCCCCCCA CATOCCCCCCA CATOCCCCCCA CATOCCCCCCA CATOCCCCCCA CATOCCCCCCA CATOCCCCCCA CATOCCCCCCA CATOCCCCCCCA CATOCCCCCCCA CATOCCCCCCCA CATOCCCCCCCA CATOCCCCCCA CATOCCCCCCCA CATOCCCCCCCA CATOCCCCCCA CATOCCCCCCA CATOCCCCCCCA CATOCCCCCCCA CATOCCCCCCCA CATOCCCCCCCA CATOCCCCCCCA CATOCCCCCCA CATOCCCCCCCA CATOCCCCCCCA CATOCCCCCCCA CATOCCCCCCCA CATOCCCCCCA CATOCCCCCCCA CATOCCCCCCCCA CATOCCCCCCCCA CATOCCCCCCCCA CATOCCCCCCCCCC		(C) STRANDERNESS: single	
(ii) MOLECULE TYPE: DEA (genomic) (xi) SEQUENCE DESCRIPTION: SEQ ID NO:6: (xii) SEQUENCE DESCRIPTION: SEQ ID NO:6: OUTCOCCOGA TOPOGROCO A CREAT COME CONTINUEDA CARCETORIA: ANDIGOROCO DOCTAGALACIA CERCONOCAM ATMANICADO CONTINUEDA CARCETORIA: ANDIGOROCO DOCTAGALACIA CERCONOCAM ATMANICADO CONTINUEDA CARCETORIA: ANDIGOROCO DOCTAGA COMENCIA CONTINUEDA CARCETORIA: ANDIGOROCO CARCETORIA: OUTCOARCE ATMANICADO CONTINUEDA CARCETORIA: ANDIGOROCA CARCETORIA: OUTCOARCE CONTINUEDA CARCETORIA: THOCKOMO CONTINUE ATMANICA CONTINUEDA CONTINUEDA TOTORISMO CONTINUEDA CARCETORIA: OUTCOARCE CONTINUEDA TOTORISMO CONTINUEDA CARCETORICO CONTINUEDA CARCETORICO GORGADA CONTINUEDA CARCETORICO CONTINUEDA CARCETORICO GORGADA CONTINUEDA CARCETORICO CONTINUEDA CARCETORICO COCOGADA CONTINUEDA CARCETORICO CONTINUEDA CARCETORICO COCOGADA CONTINUEDA CARCETORICO CONTINUEDA CARCETORICO COCOGORIA CARCETORICO CONTINUEDA CARCETORICO CONTINUEDA CARCETORICO COCOGORIA CARCETORICO CONTINUEDA CARCETORICO COCOGORIA CARCETORICO CONTINUEDA CARCETORICO COCOGORIA CARCETORICO CONTINUEDA CARCETORICO COCOGORIA CONTINUEDA CARCETORICO CONTINUEDA CARCETORICO CONTINUEDA CARCETORICO CONTINUEDA CARCETORICO COCOGORIA CARCETORICO CONTINUEDA C		(D) TOPOLOGY: Linear	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:6: 0007000003 TOURS COCCEARAGE ATTROCUSTS CONGENCIAGE TOURS CONTROLL 3AGUCARCT DECTAGRACE CARGOCCAS ATGARTAGES CONTROLLA CARGOTORIC 3AGUCARCT DECTAGRACE CARGOCCAS ATGARTAGES CONTROLLA CARGOTORIC CARGOCTCCA ACCAGINGO CARGOCCAS ATGARTAGES CARGOTORICA CARGOTO	2	(11) MACH YOUNG TO HOLYON DOWN (A DAMAGE)	
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15 GUGCCATCHAT CUGGGCTCAN GUGGGGTTGC TORROGOGNA GUATCAGGCC ATCATTOCHTO AGGGCTGAC GUGGGGTAA TOTACHAGAA AGCCAACAGC GUGGGGCAGA TTACLCAGTT GGGCCGTAAC TTCCAGGTGA TCTACCACAGC GUGGGGCAGA AGUTGCAGACA TCCAGGTGA TCTACCACAGC GUGGGGCAGA CUTTACCCAGC GGCCAGGCC AGCACACACACCAGC GUGGGGCAGA CUTTACCCAGCAGCC AGCCAGGCC AGCACACACCAGC GUGGGCAGACACCACCACCACCACCACCACCACCACCACCAC			300
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AGUTGORAGO TOCOGECANE ARCATOSCOP ANACOGRAGA GUCCOTENSO TOCAGOTOGO 54 CUTARCACCA GUCCAGAGOCA ROBADOTOS TOMENSANTO AMBOTOCO 50 TOCOGRIGOCA GUCCAGAGOCA TURACTORIA GUTTACCACCA GUCCAGAGOCA GUTTACACCA GUTTACCACACACACACACACACACACACACACACACACAC			480
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SGCCCGTTCS GGGANARCCC CARGAGGTC GGTTCAGGCT GTTACTCTCT GGTTGAACGC TTCCTGGGGG CCACCGCTTG GGGCCTGAGGCT TTTTGCCGCC ATTCCTCGGGCT GGTTGAACGC GGTGGCCAAC GACTATGACT AGGACACCGT TTTTGCCAGG GCCCTCAAA GGACTCCGGC CCCGCAAAAC GCACACCTTTTT GCGGATAAGGACACCTTTTTGCCAGG GCCCTCAAAA GGACTCCGGC CCCGCGAAAAC GCACACCTTTTTACTGAGGATCAGCAAAACACCTTTTGCCGGAAAAC GCACACCTTTTTATCAT GGGATCTCCGGAAAAC GCACACCGGAAAAC GCACACCTTTTATCAT GGGATCTCCGGAAAACACCGGAAAACACCACCTTTTATCAT GGGATCTCCGGAAAACACGGAAAACACCGGAAAACACCACCTTTTATCAT GGGATCTCCGAAAACACCGGAAAACACCGCGAAAACACCACCACCAC			780
THOLTOGOGO CCACCOCTTO GUGCCHANGO ACTECACOCC ANTOCOTONA GARTACAGO 50			
25 ggragocaac gactatuact addacaccoff titisgocad goccitonaa guarcidodo coccodaanac docaadguad tigogotsia cogurentuc aartaloogia guarcidodo coccodaanac docaadguad tigogotsia cogurentuc aartaloogia guarcidodo coccodaanac docaadguad tigogotsia cogurentuc aartaloogia guarcidodo ggcomactigo engalantoca castitetti attentica guarcidoc guarcidodo guarcidodo attentica titistituda guarcidot guarcidodo guarcidodo attentica titistituda guarcido aartaloogia guarcidodo attentica titistituda guarcidodo aartaloogia guarcidodo attentica caccaatigot gretoticot guarcidod aartaloogia aatoloogia attentica caccaatigot gretoticot guarcidod aartaloogia guarcidodo aartaloogia guarcid			
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35 OCTIGICATICE GARGIGIPOE TOATAGATHOA ATHICAGUET TOGECOCACE ACCONTOGICA COCOTOTOGIC TOCCUTARGE CORRACTOR ATHICAGUET TOGECOCACE ACCONTOGICA COCACACTUMA CCATTOCTOR BEATCOTIST TOGGGARGAGE COLORACTOR CACACAACOT COLORACTOR CACACAACOT DOLOGICATE TOCCUTARON TOGGGARGAGE TERRICATIGG CACACAACOT DOLOGICATE TOCCUTARON			
GOODTOTORO TRACETARIAS CORRACCOGO TUTASTOCTO GARTHACOCA GARCALTACO CORRATTAM CONTROCTA DESTRUCTOR TOSGERERGE CREACAGNT TENENTEGRA CARLARCET DUCUBUTET GEOGTOCAT GTGCCAGTGT CERRATAGGG GGCCCAGTT CTGCCGTGT COCTARCOCA TENTTUNCT CONTROCAGTG GTGCCAGTGT CERRATAGGG GCCCAGTT 40 ATSCEPENGG GARCAGGG TATOTOCAGT COTTCCGGG GTGCCCCT MCCGGTTCGCA ATSCEPENGG GARCAGGG TATOTOCAGT CTGCCAGC CAGCCCT MCCGGTTCGC (2) ENFORMATION FOR SEQ ID NO:7: (1) REQUENCE CHARACTERISTICS: (A) LENGTH: 2921 base pairs (B) TYPE: mucleic acid (C) STRAMEBENERS: single (D) TOPOLOGY: Binear (ii) MULBCULE TYPE: DNA (genomic) (xi) SEQUENCE CHARACTETISTICS: (xi) ESQUENCE CHARACTETISTICS: (xi) ESQUENCE CHARACTERISTICS: (xi) ESQUENCE CHARACTERISTICS: (xi) ESQUENCE CHARACTERISTICS: (xi) SEQUENCE CHARACTERISTICS: (xi) GROUNCE CHARACTERISTICS: (xi) GROUNC	35		
CONCRITING CCATTOCTON BEATCOTOST TOGGERIAGE CTENSCORT TRANSATISS CACACARACT DOLOGERICS ESCURIOR TOGGERIAGE CTENSCORT TRANSATISS CACCARACTO DOLOGERICS TOTOTOGRA DECORTOST CAMBRIDGE GOOGGERIAGE TOGGERIAGE COCCAGACUCE TRATTORIC COCCAGO CABACCOTO DELAFICURA ATUGERIAGE COCCAGACUCE TRATTORIC COCCAGO CABACCOTO MOCOGRICOS 180 (2) EMPORMATION POR SEQ ID NO17: (1) SEQUENCE CHARACTERISTICS: (A) LEMOTH: 2921 base pairs (B) Type: mulcic acid (C) STRAMBENESS: siegle (D) TODOLOGY: 318-AX (11) MULBOCUE TYPE: DNA (GENOMIC) (X1) SEQUENCE DESCRIPTION: SEQ ID NO17: 55 COGURISCO TRATGETT CATTOCCAA ACCOTOGOR TROTCCUCOG GGTATCCAGG TCCGGOTOR CCATCAGOGC TRATGETT COTOGRATIC ACCOTOARCT GGCOGCCCAA 12 TCCGGOTOR CCATCAGOGC TRATGETT COTOGRATIC ACCOTOARCT GGCOGCCCAA 12 TCCGGOTOR CCATCAGOGC TRATGETT COTOGRATIC ACCOTOARCT GGCOGCCCAA 12 TCCGGOTOR CCATCAGOGC TRATGECT TOTOGGATIC ACCOTOARCT GGCOGCCCAA 12	24		
CACACARCET DECORTECT ECCUSTORM TSTOCCASTST COMMITTAGES GOCGOCAST 124 CTGGCGGTC COCTAGEACCE TENTION CONCINCION ACTORISTO COMMITTAGE CONCINCION ACTORISTORM COMMITTAGE CONCINCION COMMITTAGE CONCINCION COMMITTAGE CONTROL 126 ACTEMICACET SECURION FOR SEQ ID NO.7: (1) EMPORENTION FOR SEQ ID NO.7: (A) LENGTH, 2921 base pairs (B) TYPE: mucleic acid (C) STRANDEDNESS single (D) TOPOLOGY: linear (ii) MULBOULE TYPE: DNA (genomic) (xi) SEQUENCE DESCRIPTION: SEQ ID NO.7: 55 COMMITTAGE TRANSPORTES TO TORIGATION ACCORDANCE GOCGOCCOS TOCCOGNACT GOCGOCCOS TOCCOCCOS TOCCOCCOCCOS TOCCOCCOCCOS TOCCOCCOCCOS TOCCOCCOCCOS TOCCOCCOCCOS TOCCOCCOCCOS TOCCOCCOCCOS TOCCOCCOCCOS TOCCOCCOCCOS TOCCOCCOCCOCCOS TOCCOCCOCCOS TOCCOCCOCCOCCOS TOCCOCCOCCOCCOCCOS TOCCOCCOCCOCCOS TOCCOCCOCCOCCOCCOCCOCCOS TOCCOCCOCCOCCOCCOCCOCCOCCOCCOCCOCCOCCOCC			
CTCCCCCTCCC CCCCCCCCCCCCCCCCCCCCCCCCCC			
40 AFGGREGGG GGAGCGGGG TRATEGCGAT CTTGCGCGGG GGAGCGCGT MGCGGNTCGG 186 ACTMMGCGGT GGGGGACAG ACGTGGACC CTACTCGGGC CAGTT (2) INFORMATION FOR SEQ ID NO:7: 45 (1) SEQUENCE CHARACTERISTICS: (A) LEMOTH: 2921 base pairs (B) TYPE: mucleic acid (c) STRADEBONES: sidgle (D) TOFOLOGY: lineax (ii) MULBCGLE TYPE: DNA (genomic) (xi) SEQUENCE DESCRIPTION: SEQ ID NO:7: 55 CDGGGTGGA CCATCAGGGC TRATEGCGAA ACCGTGGACT GGCGGCGGA TCGGGGTGA CCATCAGGGC TRATEGCTCT TOTGGATTCH ACGGTGAACT GGCGGCGGA TCGGGGTGA CCATCAGGGC TRATEGCGTGT TCGGGGTGCT GGCGGCGGACT GGCGGCGGACT TGGCGCGCGACT GGCGGCGGACT GGCGCGCGGACT GGCGGCGGACT GGCGCGGACCGGAC			
ACTHRECOGT GGOGGACAG ACETGGACC STACTCGAGC CAGTT (2) EMPORMATION FOR SEQ ID NO:7: (1) SEQUENCE CHARACTERISTICS: (A) LEMOTH: 2921 base pairs (B) TYPE: mucleic acid (C) STRANBENESS: stopie (D) TUDOLOCY: 318-ex (ii) MULBOULE TYPE: DNA (genomic) (xi) SEQUENCE DESCRIPTION: SEQ ID NO:7: 55 CDGUARGCOS TEGUENTES TRANSCOCAA ACCOTOGOGC TEGUENCOCAC GGCAGCCOGAC TEGUENCOCAC 12 TCCGGGTGAC CCATCAGGGC TUGACTGGTT COGGGTGACT GGCCGGCCGA TEGUENCOCAC 12	435		
(2) INFORMATION FOR SEQ ID NO:7: 45 (1) SEQUENCE CHARACTERISTICS: (A) LEMOTH: 2911 base pairs (B) TYPE: mucleic acid (C) STRAMBEDNESS: single (D) TOPOLOGY: linear (ii) MULRICULE TYPE: DNA (genomic) (xi) SEQUENCE DESCRIPTION: SEQ ID NO:7: 55 COGGREGO TEGRUENTED TATTSCOCAA ACCOTOGOGY TROTCOCCOG GGTATCCRGG TCCGGGTGGA CCATCAGGGG TRUBLETGTT TOTGGATTCH ACCGTGAACT GGCCGGCGGA TTCCGGGTTTCAGGATTCT NOCAGGGTT TTCGGGTTCTC GTTTCCCGAC 12	**()		
45 (1) SEQUENCE CHARACTERISTICS: (A) LEMOTH: 2921 base pairs (B) TYPE: mucleic acid (c) STRANDEDNESS: single (D) TOPOLOGY; linear (ii) MOLECULE TYPE: DNA (genomic) (xi) SEQUENCE DESCRIPTION: SEQ ID NO:7: 55 CEGUATGECE TEGROUTTEG TATTSCOCAA ACCOTOGOGY TOGOCOCOG GGTATCCRGG TCCGGGTTGA CCATCAGGG TRUBACTGGTT TOGGGATTCG ACCGTGAACT GGCCGGCGAAC TTCGGGTTGA CCATCAGGGG TRUBACTGGTT TOGGGATTCG ACCGTGGAACT GGCCGCCGAACT		sectionactions: and characteristic versitations of assessment assessment consist	2302
(A) LEMOTH: 2921 base pairs (B) TYPE: nucleic acid (C) STRANDEDWESS: single (D) TOPOLOGY: linear (ii) MOLDCULE TYPE: DNA (genomic) (xi) SEQUENCE DESCRIPTION: SEQ ID NO:7: 55 COMMINGCOS TROTOGYTES TRUBENGETT TORGENICO ACCGROACT GEOCOGCOS TCCOGOTOGA CONTEAGOGY TROGENICOTOGO GETATOCOGA 12 TCCOGOTOGA CONTEAGOGY TROGENICOTOGO GETATOCOGA 12 TCCOGOTOGA CONTEAGOGY TROGENICOTOGO GETATOCOGAC 18		(2) INFORMATION FOR SEQ ID NO:7:	
(A) LEMOTH: 2921 base pairs (B) TYPE: nucleic acid (C) STRANDEDWESS: single (D) TOPOLOGY: linear (ii) MOLDCULE TYPE: DNA (genomic) (xi) SEQUENCE DESCRIPTION: SEQ ID NO:7: 55 COMMINGCOS TROTOGYTES TRUBENGETT TORGENICO ACCGROACT GEOCOGCOS TCCOGOTOGA CONTEAGOGY TROGENICOTOGO GETATOCOGA 12 TCCOGOTOGA CONTEAGOGY TROGENICOTOGO GETATOCOGA 12 TCCOGOTOGA CONTEAGOGY TROGENICOTOGO GETATOCOGAC 18			
(B) TYPE: nucleic acid (C) STRANDENDERS: single (D) TOPOLOGY: Binear (ii) MULBOULE TYPE: DNA (genomic) (xi) SEQUENCE DESCRIPTION: SEQ ID NO:7: 55 COGGREGO: TOGGREGORY TOGGREGORY TOGGRACT GEOCOGCOR 12 TCCGGGTCGA CCATCAGOGC TUGACTGTT TOTGGATTCG ACCGTGAACT GEOCOGCORA 12 TCCGGGTTCG TCGGCGCAT TOGGCGTT TOGGGCTCCOGGCT TOGGCCGCAC 1	45		
(C) STRANBENNESS: single (D) TOPOLOGY: linear (ii) MULHCULE TYPE: DNA (genomic) (xi) SEQUENCE DESCRIPTION: SEQ ID NO:7: 55 COGGREGO: TOPOGRETIC TATTSCOCAA ACCOTOGOCO TOGOCOCOA GETATCORGE TOCOGOTOGA COATCAGOGO TROACTOTT CORGARITOC MACGOTOGAC TOGOCOCOAC 12 TOCOGOTOGA COATCAGOGO TROACTOTT CORGARITOC MACGOTOGAC			
(D) TOPOLOGY: linear (ii) MULBOULE TYPE: DNA (genomic) (xi) SEQUENCE DESCRIPTION: SEQ ID NO:7: 55 COGULTECCO TGGTGGTTGG TATTSCCCAA ACCCTGGCGC TGGTCCCCGG GGTATCCRGG E TCCGGGTGG CCATCAGCGC TUGACTGTT TCTGGATTCG ACCGTGAACT GGCCGCCGAAC TGCCGCCGAC TGGCCGCCGCAC TGGCCGCCGAC TGGCCGCCGAC TGGCCGCCGAC TGGCCGCACC TGGCCGCACC TGGCCGCACC TGGCCGCACC TGGCCGCACC TGCCGCACC TGCCGCACC TGCCGCACC TGGCCGCACC TGCCGCACC TGCCGCACC TGCCGCACC TGCCGCACC TGCCGCACC TGCCGCCACC TGCCGCACC TGCCGCACC TGCCGCCACC TGCCGCCACC TGCCGCCACC TGCCCCCACC TCCCCCCACC TCCCCCCACC TCCCCCACCCA			
50 (ii) MULRCULE TYPE: DNA (genomic) (xi) SEQUENCE DESCRIPTION: SEQ ID NO:7: 55 CEGUARGECT TRATESPECTE TATESPECAA ACCOMPAGE TRATESPECT TOGGATTCA CONTEASON TRASECTORY TOTOGGATTCA SECURIORIST TOTOGGATTCA SECURIORIST TRASECTORY TRASECTORY TRASECTORY TRANSPORT TOTOGGATTCA SETURICAL 18			
(ii) MULRCULE TYPE: DNA (genomic) (xi) SEQUENCE DESCRIPTION: SEQ ID NO:7: 55 COGGREGO TEGROGETEG TATTSCOCAA ACCOTOGOGO TEGROCUCOG GGTATCORGG E TCCGGGTGG COATCAGGG TEGROGETEG TAGGGTGT TCGGGATCT TAGGGTGT TATTSCOCAC 18	**	(D) TOPOLOGY: linear	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:7: 55 COMMUNICOS TRATEGORA ACCOTOGOGO TEGROCOCOS GETATOCADE E TOCOGOTOGA COATCAGOGO TEGRACOTOTO ACCOTOGACTO GEOGOGOCOA 12 TOCOGATOCO TRATEGRACOTOT TROGOCOTOCO SETUPITETE SPERIOCOGAC 12	50	The second secon	
55 COGGREGOS TGGTGUTTGG TATTGCCCAA ACCOTOGOGO TGGTGCCCGG GGTATCCRGG CCGGGGTCGA CCATCAGOGO TGGACTGTT CYCGGACTGG ACCGTGAACT GGCGGCCCGA TCGGGGTCC TCGGGGTCCC GGTTGTTCTC GTTGCCCGAC 12		(ii) NOLECULE TYPE: DNA (genomic)	
TCCGGGTCRA CCATCAGORC TUGACTGTFT CYCGGACTCG ACCGTGRACT GGCCGCCGA 12 TTCGGRTTCC TGCTGGCCAT TCCAGCGGTG TTCGCCTCCG GGTTGTTCTC GTTGCCCGAC 18		(xi) SEQUENCE DESCRIPTION: SEQ ID NO:7:	
TCCGGGTCRA CCATCAGORC TUGACTGTFT CYCGGACTCG ACCGTGRACT GGCCGCCGA 12 TTCGGRTTCC TGCTGGCCAT TCCAGCGGTG TTCGCCTCCG GGTTGTTCTC GTTGCCCGAC 18	55	CONTAINATION BYSTHANIAGA WEIGHNOON A MANAGONO ARRIVATOR AND ARRIVATOR	
TYCGGATYCC TGCTGGCCAT TCCAGCGGGG TYCGCCTCCG GGTTGTYCTC GYTYCCCGAC 18	272		
perimente postantia dacatame perimente estantino, dalacete. 31			
		GUITECATE PROMOCTOR GOOGRIGHT RELEGISCE CACARLIBOL COLOROGE	241

	CTGATCGCGT	TEGTECTEGG	TCTGACCGCG	GYGGCCTGGC	TOCTOCOGTE	TCTGOTGCGA	300
	CACAACATGT	ACTOGTTCOT	CGGCTACCGG	@1/@CTO9TO3	GGACGGGCAT	SCTCSTSCTS	360
	CYGGCTACOG	GGACGGTAGC	CCCGACATGA	CCGTCATCTT	GCTACGCCAT	SCCCGTTCCA	420
	CCTCGAACAC	cocogocoro	CTOGCCGGCC	COTCOMMONT	CGACCTCGAC	GAGAAGGGGC	480
- 5	GCGAGCAGGC	CACCGGGTTG	ATCGATCGAA	TIGGIGACCT	OCCUATOOSG	GCGGTCGCGT	340
	CFICTCCAAT	GCTGCGGTGT	CRACGCACCG	TOSSANCOSCT	GGCCGAGGCG	characetas	500
		COATGACCOG					660
	TOGGTGACCT	GOTCGACGAG	CCGTTGTGGC	GGGTAGTCCA.	GGCCCACCCC	AGOGGGGGGG	720
	TOTTTTTCCCCOC	COGTGAGGGT	TYGGCGCAGG	TECAGACETS	GTTGTCCTGA	COGATITCCA	780
36	TOCCOMPGAA	CACCAAGACC	GGATOGGCAC	TEGGETTEGE	COGCONANAC	CCGGCCCGCCA	840
-	ATAGGGGGAC	COTTOSCIGOS	AATGCGCGTG	GTACCA6GOG	SACCACCTTG	AACTOCCATC	900
	CGTCGGGGCC	AAGCGCATCG	ccccccccc	OPTACEGOTA.	AGGCOTACCA	AAACCCGACG	960
	GTANTACTTC	GOCAANOTOO	GOTTONOGACG	TTACOGAGAC	GTGACCAGNG	AGGCNGCGGC	1.020
	ATTGGATTTA	TOGATGOTGC	SCHOTTCOCA	NCCCGGCGGT	COGAANACGT	AGCCCAGCCG	1060
3.5	ATCCCGCAGA	COTOTTOCCO	ACCGCCAGTC	ACGCACGATC	COCACGTACT	COCGGGGTCTO	1140
	CASCTTCCAG	ATGTTGAACG	TOTOGROUG	CTTGGTCAGO	CCATAATGCG	GTCGGAATAG	1200
	CTCCGGCTGA	AAGCTACOGA	ACAGGGGGTC	COAGATGATG	AGGATGCCGC	CATAGTTCTT	1260
	- GTCCANATAC	ACCOGGGTCCA	TTCCGTGGTG	GACCCGGTGG	TGCGACGGGG	TATTGAAGAC	1320
	CARCOCCARO	CACCGCGGGCA	GCCTGTCGAT	ccccrosers	TOCACCCAGA	ACTOGRADAT	1300
20		CACCAATTGC					1440
	AACCCACATG	AGAATCTCGC	CCCTCTTCTT	CCAMPTTCTG	OCGCAGCGCG	GTGGCGAAGT	1500
	TGAAGTATTC	GCTQQAQTQA	TOCGCCTOOF	OGGTAGOCCA	GATCAGCCGA	ACTCGGTGGG	1560
	CGATGCGGTG	ATAGGAGTAG	TACAGCAGAT	CGACACCAAC	GATOGOGATO	ACCCAGGIGT	1620
	ACCACCGGTG	GGCGGACAGC	TOCCAGGGGG	CAAGGTAGGC	ATAGATTGCG	GCATAACCGA	1680
25	GCAGGGCAAG	GGACTTCCAG	CCGGCGGTGG	TUGCTATCGA	AACCAGCCCC	ATCGAGATGC	1740
	TGGCCACCGA	GTCGCGGGTG	ACCTAACCCC	CCGAGGCGGG	CCGTGGCTGC	CCCGTAGCAG	3,800
	CHETCTOGAT	GCTTTCCAGC	TTGCGGGGGG	CCCTCCATTC	GAGAATCAGC	RGCRATAGAA	1860
	AACATOGAAT	GGCGAACAGT	ACCUGUTCCC	GCRTTTOCTC	GGGCAGCGCT	GAGAAGAATC	1920
	COGCOACOOC	ATOGCCGAGG	COACCTCOMT	AGACACCATO	ACCCAGATGA	ATCAGGGGTT	1980
30) TOGGARGATO	GIGAACATGC	TGCACGGGGT	GCGTGACGGG	Chachacaca	ACGCCAACAA	2040
	NTACGAACAG	CAAGAGCAGG	CCTCCCAGCA	CATCCTCAGC	AGCTGACCCG	GCCCGACGAC	2100
	TCAGGAGGAC	ACATGACCAT	CAACTATCAA	TTCGGGGGACG	TOGACGCTCA	COOCOCCATG	2160
	ATCCCCCCCTC	AGGCCGGGTC	GCTGGAGGCC	CAGCATCAGG	CCATCATTTC	TUATUTUTU	2230
		ACTTITGGGG	CGGCGCCGGT	TCGGCGGCCT	OCCAGGGGTT	CATTACCCAG	2280
35	CTUGGCCGTA	ACTTCCAGGT	GATESTACGAG	CAGGCCAACG	CCCACGGGCA	GAAGGTGCAG	2340
		ACAACATOOC					2400
	TOGCTTAAGG	cccacaccon	CAATTACAAC	OTGGCCGCAC	ACCOGTTOGT	GTGTGGCCAC	2460
		GAACGACTAA					2526
	CCATGGTGCT	GAACTGGGAA	GATOGCCTCA	RIGCCCTTGT	TOCCOANOGC	ATTGAGGCCA	2580
4() TOGTOTITOS	TACTITAGGC	GATCACTCCT	COTTOTOGGA	GICGCIGCIG	CCCGACGAGG	2640
	TOCCCCCACT	GCCCGAGGAA	CTGOCCCGGG	TGGAGGCATT	GTTGGACGAT	CCGGCGFFCT	2760
	TOSCOCOMT	COTOCCOTTC	TTOGACCCGC	GCAGGGGCCG	GCCGTCGACG	CCGATGGÁGG	2760
		GTTGATGTTT					2620
		TGATTCGATC				CACCICTOGC	2880
45	TOCCOCATOC	GACCACATTG	ATGRAGOTCA	CCYCGCGLLA	C		2921

(2) INFORMATION FOR SEQ ID NO:8:

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- (i) SEQUENCE CHARACTERISTICS:
 - (A) LEMOTH: 1704 base pairs (B) TYPE: nucleic acid
 - (C) STRANDEDMESS: single
 - (D) TOPOLOGY: linear
- 55 (ii) MOLECULE TYPE: DNA (genomic)
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:8: